

CLINICAL ARRAY READER (CAR) USER MANUAL



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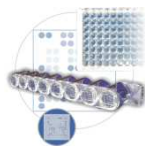

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2. INTRODUCTION

The *Clinical Array Reader (CAR)* has been designed for a quick detection and analysis of ASs. This detection and analysis represent the second step of an CSs assay. The first step of a complete CSs assay also involves experimental procedures (sample preparation, DNA extraction, DNA amplification and target labelling, specific hybridisation and visualization of the results).

The detection and analysis of CSs by the *CAR* is always performed immediately after developing the specific precipitation staining pattern on the CSs. Developing of this pattern generates a dot micromatrix (microarray) on the t the bottom of the CSs. This micromatrix can be detected and analysed by the *CAR*. The results obtained by the *CAR* are automatically shown on the touch screen and can be printed or exported to your LIMS or pen-drive by its USB connections.

Experimental Procedures	Sample preparation, DNA extraction and DNA amplification/labelling of targeted molecules	Standard laboratory equipment
	Specific hybridisation: labelled sample incubation against arrays. Visualization: conjugation and precipitation of the staining.	
Visualization	Colorimetric Detection and Analysis.	 CAR (Clinical Array Reader)

3. COMPONENTS DESCRIPTION



Performance:

Resolution on the sample: 3 μm
 Camera pixel resolution: 8.3 Mpixel
 Scan time for complete 96-well plate: 2-3 minutes
 Dynamic Range: 8-bits, 255 greyscale

Mechanical:

<i>Dimensions</i>		
<i>Width</i>	<i>Height</i>	<i>Depth</i>
44 cm	21 cm	34 cm

Additional **10 cm** space must be provided in front of the instrument for microplate loading, as well as an additional **10 cm** space at the back for cables and ventilation.

Weight (without tactile screen) **15 Kg**
 Replacement fan filter 80x80mm NMB, Part No.: F80/MR

Environmental:

<i>Environmental Conditions</i>		
<i>Operating Temperature</i>	<i>Storage Temperature</i>	<i>Relative Humidity (non-condensing)</i>
0 – 70 °C	-40 a + 85 °C	10 – 75 %

Electrical:

Electrical Specifications		
Rated Power	Fuse type and rating two fuses	Voltage
150 W	5x20 mm, 250V 1.6A slow-blow	110 – 230 V (AC) a 50/60 Hz

Image storage format 8-bit BMP greyscale

Results storage format CSV spreadsheet

Internal PC:

Specifications	
Operating System	Windows XP home edition
Processor	Intel Pentium II 1.6 GHz (o mayor)
RAM Memory	1 GByte DDR-2
Hard Disk	40 GByte
Connections	VGA, RS-232 (mouse and keyboard), Serial100 Base-T Ethernet, 3x USB2

Touch screen:

Characteristics	
Type	12.1" TFT LCD
Resolution	800x600
Colours	264 K
Connections	12V, VGA, USB-B,

Mechanical mount VESA 75

Max current for 12VDC tactile screen supply: 3 Amp.

4. INSTALLATION



Please, follow the indications described in this manual to set up the Clinical Array Reader (CAR).

4.1 PRECAUTIONS

- Verify that the instrument does not present any obvious sign of damage. **If any damage is found, please contact immediately to your local GENOMICA's representative.**
- Prepare approximately 1 m³ free area in the lab for the installation of the CAR. Once unpacked, place your CAR in a standing position on the bench space you have just prepared. Relocation of your CAR is only recommended by lifting it, **DO NEVER PUSH YOUR CAR.**
- Make sure that the CAR is placed on a stable and perfectly horizontal bench.
- Before connecting it to the power supply, make sure that your power line voltage corresponds to the **operating voltage of 110-230 V (AC) at 50/60 Hz.**
- Do not install the CAR in the influence area of a heat or intense light source.

4.2 UNLOCK/LOCK THE READER



In order to avoid system damages during transport, internal mobile components of the CAR are locked. In order to unlock the reader, remove the big screw on the rear side and on the right side.

Readers contain two locking systems, one on the rear side and one in the right side.

During transportations, the position of the tray holder is fixed by lock screws, in order to protect the mechanical components. Remove these screws before switching on the instrument for the first time and keep them for the posterior assembling of the screen. No special tools are required to remove both screws.

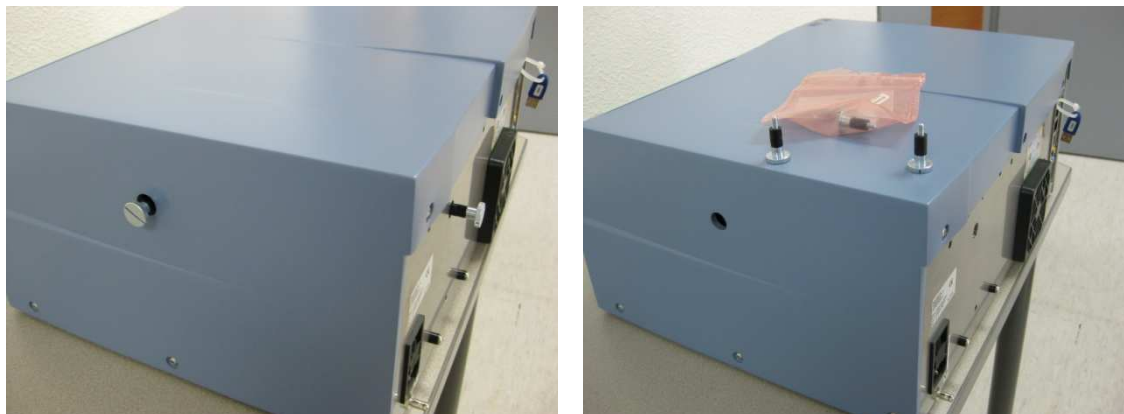


Fig.1: Unlocking the CAR. Position of the locking screws.

Place the white plastic cap attached in the hole left by the lock screw at the side of the CAR (Fig. 2)



Fig. 2: Covering the side hole left by the screw..

Should you need to send your instrument, the lock screws **must** be screwed securely in their positions. Unscrewed readers are susceptible of further damages during transportation.

To install the transport lock screws, please follow these short instructions:

- Turn off the instrument and remove the main power connection.
- Slowly slide the tray holder manually fully towards the rear wall and place the screw at the rear of the CAR.
- Then slide the tray holder fully to the right and place the screw on the side.
- Tight them securely.

4.3 ASSEMBLING THE SCREEN

Touch screen is provided separately. In order to connect it, please follow these indications:

1. Unscrew partially the two lower screws (yellow), and totally those marked in red, all of them are placed on the rear side of the reader.



2. Lean the screen over the reader fitting the screws with the positions marked in yellow. Insert the safety lock screws recently removed (green circles, below) in the positions marked in red (See below).



3. Once installed, screen's arm should appears as follow:



4.4 CONNECTING SCREEN AND READER



1. Connecting cables:

- 1a) Connect the power supply (Type Jack 3.5)
- 1b) Connect the USB p to one port at the right side.
- 1c) Connect the VGA connection to the blue connector.

- 2. Switch on the screen by pressing the button on top.
- 3. Switch on the CAR

5. SAFETY NOTES



Please, be particularly aware of electrical hazards. Standard electrical safety precautions must be applied at all times. Only qualified personnel are permitted to perform electrical servicing.

- Never touch switchers or outlets with wet hands.
- Always switch off the instrument before disconnecting the AC power cord.
- Always unplug the instrument prior to cleaning major liquid spills and prior to servicing the electrical or internal components.
- Do not open the housing of the instrument!.
- Do not attempt any repairs or alterations except as expressly instructed in this manual or by the technical service.!
- Avoid any movement or vibration during reading operation.

6. PROCEDURAL LIMITATIONS

- *Clinical Array Reader (CAR)* has been designed for the detection and analysis of ASs/CSs (Array Strips/CLART® Strips) or ATs (Array Tubes®) only. Any other tubes, vials, or material must not be used with this equipment.
- Only use Strips or Tubes processed in accordance to the AS/CS/AT protocols provided by GENOMICA.
- Use of this equipment in a different way of that described in this manual will result in lost of your warranty rights



Please consider that the USB used for software updates or results export does not contain any antivirus. GENOMICA is not responsible about viral infections and CAR cleaning will not be considered as a part of the technical support.

7. SWITCHING ON/OFF THE CAR

7.1 SWITCHING ON

Check if your CAR requires a Dongle-key. (see FAQ 11.4). Depending on the CAR serial number, dongle-key might be already installed inside or should be installed by user. A sticker on the rear side of the CAR will inform about the position of the dongle.

Plug in the reader and switch it on. The main switcher is on the rear (right) side of the device. Once pressed, function gauge will display a yellow light. Press then the secondary switcher (Fig below), on the front side of the CAR and wait until function's gauge turns green.

Once your CAR is switched on the instrument will initialise, during this time the instrument will verify its good performance by a series of self tests. After finishing this set of self tests the "Main Menu" screen (GENOMICA application) will appear on the touch screen..



Fig.: Secondary switcher at the front side

7.2 SWITCHING OFF

First press "Shut Down" box on the main menu to switch off the CAR and then turn off the main switcher at the back side.

It is important to shut down the CAR when not required, as this may lengthen its half-life (Fig. 1).

Do not switch off the reader directly since it could damage it!!!



Shut down option at the GENOMICA'S main menu

8. READING STRIPS (AS/CSSs)



When performing CLART® Strips make sure that the metallic frame required is properly positioned on the tray, as it is shown in the image below. In case of performing Array Strips, metallic frame is not necessary.



Reading and analyzing samples is a sequential easy procedure, composed by multiple menus where user can choose among the several options of the application.

In order to start new analysis, choose option "New Analysis" on the application's main menu (Fig. 1).



Fig.2: Main menu of the GENOMICA's application

Once this option is selected, a new menu will pop up on the screen, where 96 wells microplate sketch is depicted (Fig. 3).

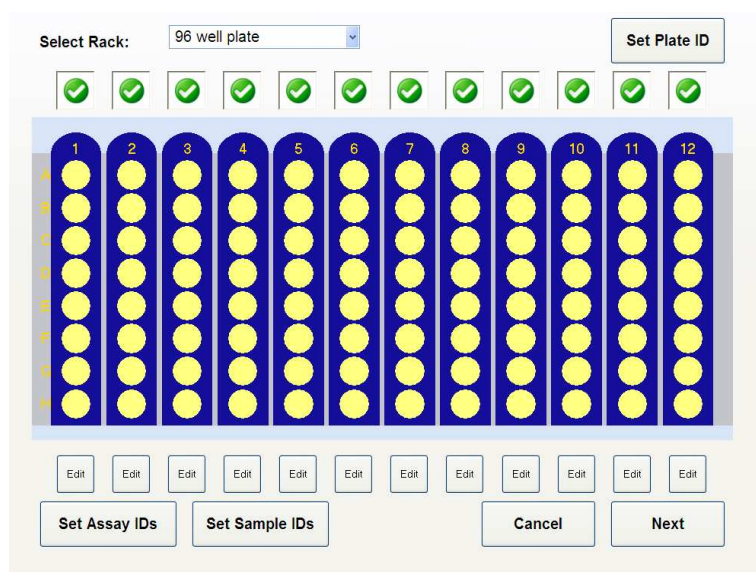


Fig. 3: 96 wells microplate sketch on the "New Analysis" menu.

This menu (Fig. 3) represents the cornerstone of the application, since it allows the user to:

- Set *rack type*: choose between 96 wells microtiter plate (AS/CS) or tubes adaptor (AT).
- Set the *sample's number* to be processed.
- Set *assay ID* (HPV2, PneumoVir, ENTHERPEX, EnteroBac+...).
- Set *sample ID* (Sample Reference).

8.1 SETTING SAMPLE'S NUMBER

The number of samples to be analyzed can be introduced manually or automatically, using a barcode reader.



It's not necessary to place the AS/CSs in correlative positions for their analysis. Nevertheless, when the CAR is included in a Clinical Array Processor (CAP), AS/CSs must be placed in correlative order, at the left side of the frame.

Automatic setting:

For the automatic setting of sample's number, user will need a bar-code reader connected to the CAR (normal USB connection)(See FAQ 2). GENOMICA can provide the user with specific bar codes for each assay upon request.

Press “Set Assay IDs” option, placed on the lower left side of the screen and all strips will turn unselected on the “New Analysis” menu (Fig. 4).

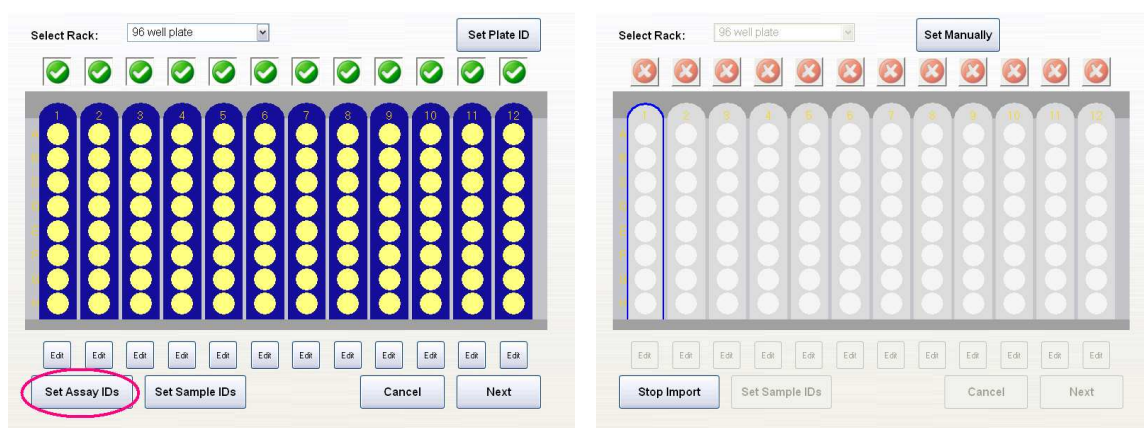


Fig.4: Left: Set Assay ID option on the lower left side of the screen. Right: All strips turn unselected when chosen this option

Every barcode read, a strip is activated, automatically, on the microplate sketch and colour combination turns from grey into a specific assay's colour combination. (Fig.5)

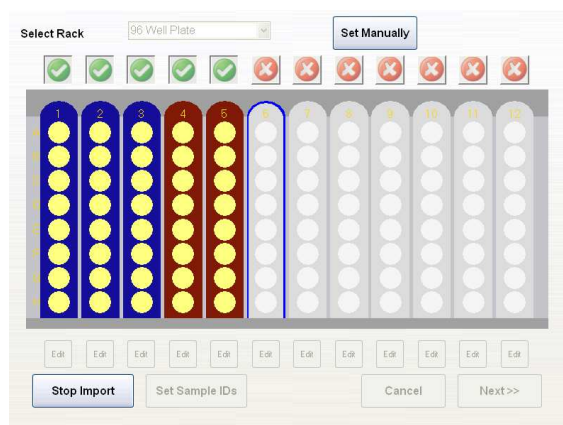



Fig. 5: Specific colours combinations. Red: HPV, Blue: Pneumovir, Green: Enterpex.

Manual setting:

Software's default settings, establish the analysis of the whole microplate, composed by up to 12 strips. Every strip not included in the process, must be manually unselected by the user.

To unselect them, proceed to press on the  symbol, which will turn into , indicating the “no selection” of the strip. The unselected strip appears on a light-grey colour (Fig. 6).

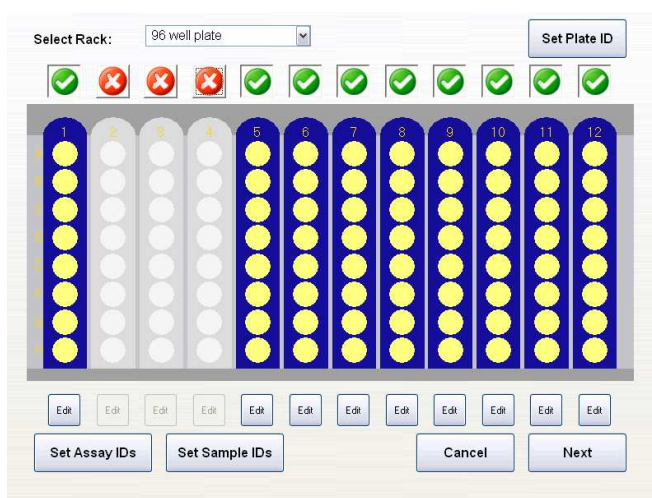


Fig. 6: Selection of AS number.




Sample's number should be, always, a whole multiple of eight, which is the number of wells included in each strip.

8.2 SETTING ASSAYS IDs

The automatic setting of sample's number, in the previous section, allows the user to skip this step.

Setting Assays IDs Manually:

Software processes by default the CLART[®] HPV2 assay. To read or analyse any other GENOMICA'S CLART[®] Assay, press the  button on the “New Analysis” menu (Fig. 7a), placed under each strip.

Option “Edit” activates a new window (Fig. 7b) that lets the user choose among those assays shown in the pull-down menu.

User can apply same Assay ID to the whole plate from the Edit menu.

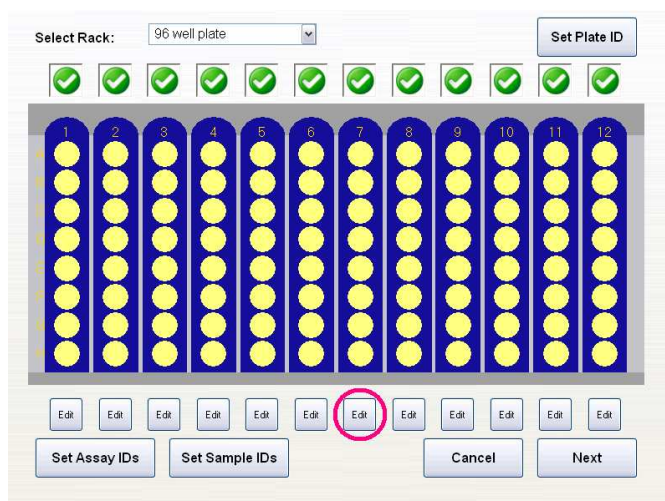


Fig.7a: Selecting “Edit” option on the “New Analysis” menu.

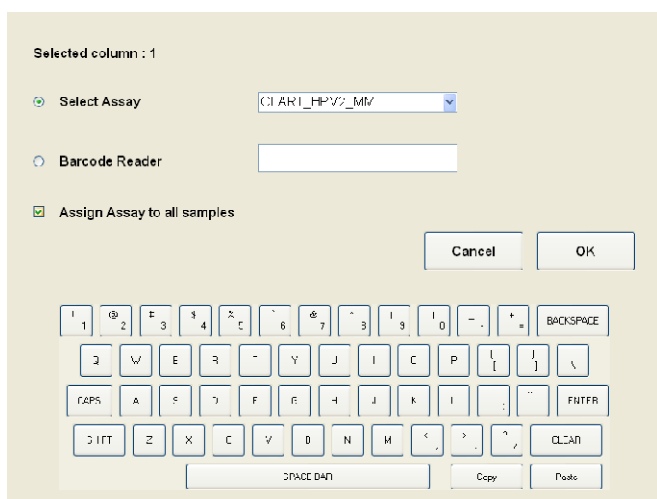


Fig. 7b: Manual setting of Assays IDs. A pull down menu shows the multiple choices to choose among

Press “OK” when finish, to return to previous menu and follow up with the visualization process.

Once assigned the assays’ IDs, strips will turn to the different colour combination specific for each type of analysis (Fig. 7C)

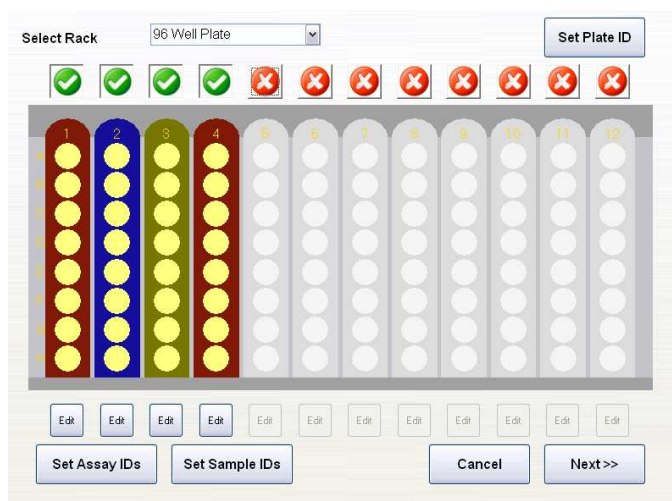


Fig.7c: Different colours combinations corresponding to different assays.

8.3 SETTING SAMPLES IDs

In the same way that in previous sections, setting samples IDs can be carried out, automatically or manually.



In case of no Samples IDs assignment, software will set automatically a correlative number to each sample, from 1 to 96, starting with the sample placed in position 1A (upper-left).

Setting Samples IDs Automatically



Barcode reader is required to set Samples IDs automatically and barcodes must be set and printed before start setting the IDs.

Set and print barcodes before starting.

Press “Set Samples IDs” option and read the previously assigned barcodes normally. The software associates automatically a barcode to a sample position. Samples with already assigned ID will appear in black (Fig. 8)

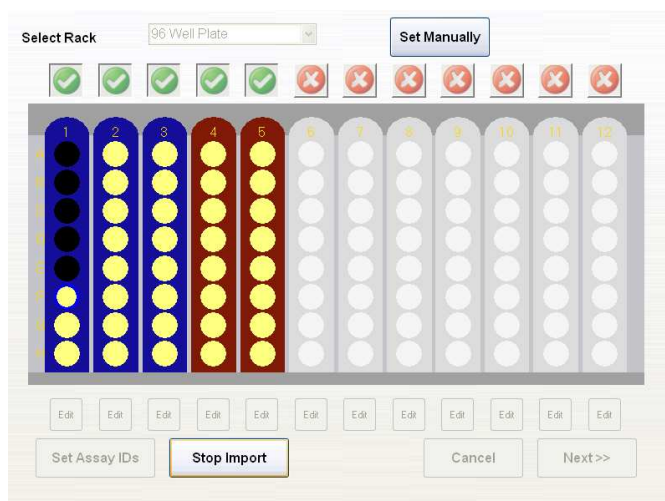


Fig 8: Setting Samples ID automatically.

Once finished importing IDs, press “Stop Import” to return to the “New Analysis” menu.

Setting Samples IDs Manually

Select each position individually by pressing twice on it. A new window representing a “tactile key board” will pop up on the screen (Fig. 9). Use it to set samples ID manually. Firstly, erase the number assigned automatically by the reader using the “Backspace” key, then insert the ID desired.

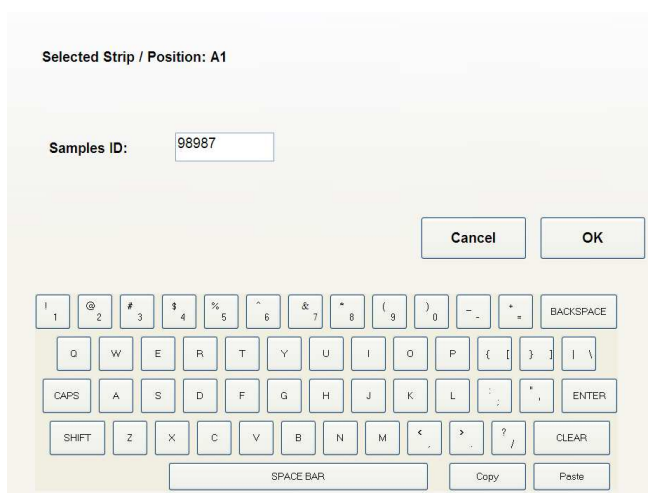


Fig 9: Setting “Samples IDs” manually, using the tactile keyboard.

The user can check the sample position at any time, since it appears at the upper-left side of the screen.

Once finished with samples’ identification process, press “OK” to go back to the “New Analysis” menu (Fig.2).

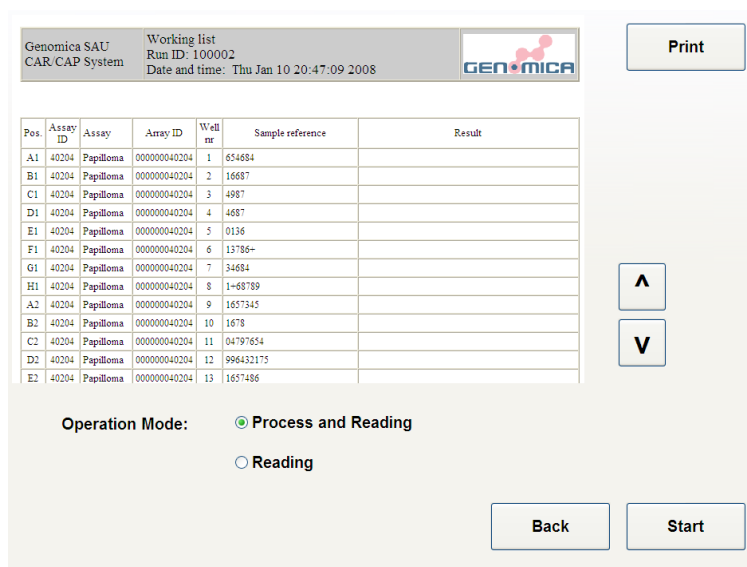
Then press “NEXT”.

8.4 WORKING LIST

The working list allows the user to check, whether the already introduced information (samples references, assays...) is correct or not.

To change any data, just press “Back” and repeat the setting process again (see “Set Samples IDs Manually” and “Set Assays ID Manually”).

If all information is correct, press “Start” to carry on with the reading process (Fig. 10).



Pos	Assay ID	Assay	Array ID	Well nr	Sample reference	Result
A1	40204	Papilloma	000000040204	1	654684	
B1	40204	Papilloma	000000040204	2	16687	
C1	40204	Papilloma	000000040204	3	4987	
D1	40204	Papilloma	000000040204	4	4687	
E1	40204	Papilloma	000000040204	5	0136	
F1	40204	Papilloma	000000040204	6	13786+	
G1	40204	Papilloma	000000040204	7	34684	
H1	40204	Papilloma	000000040204	8	1-68789	
A2	40204	Papilloma	000000040204	9	1657345	
B2	40204	Papilloma	000000040204	10	1678	
C2	40204	Papilloma	000000040204	11	04797654	
D2	40204	Papilloma	000000040204	12	996432175	
E2	40204	Papilloma	000000040204	13	1657486	

Operation Mode: ☒ Process and Reading ☐ Reading

Back Start

Fig. 10: Working list, prior to analyzing. The working list includes necessary information for the samples' traceability (references, assay ID, position on the microplate...).

8.5 READING PROCESS/ TAKING IMAGES



Reader's tray opening and closing processes are automatic. Do not try to force or move it manually. If necessary, press the emergency button placed on the rear side of the CAR.

User will be requested to initialize the process (Fig. 11), by pressing the “Open Tray” box.

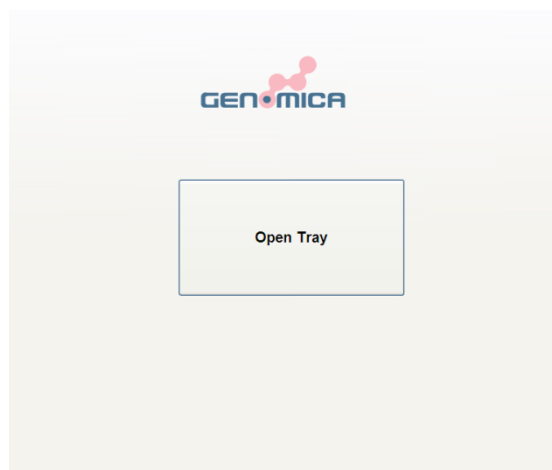


Fig.11: Message to initialize the reading process.

Once finished the opening process, place the microplate manually in the tray and operate the "Close Tray" box. (Fig.12).



Fig. 12: Message to activate the closing process of reader's tray.

Reading procedure can be monitored in the screen since samples/wells change their colour indicating the reading status (Fig.13):

- Wells in green: Sample already analysed. Grid alignment agreed.
- Wells in red: Sample may be reanalyzed. Grid alignment failed (see "Sample Reanalyze" section below).
- Wells in yellow: Sample has not been analyzed yet.

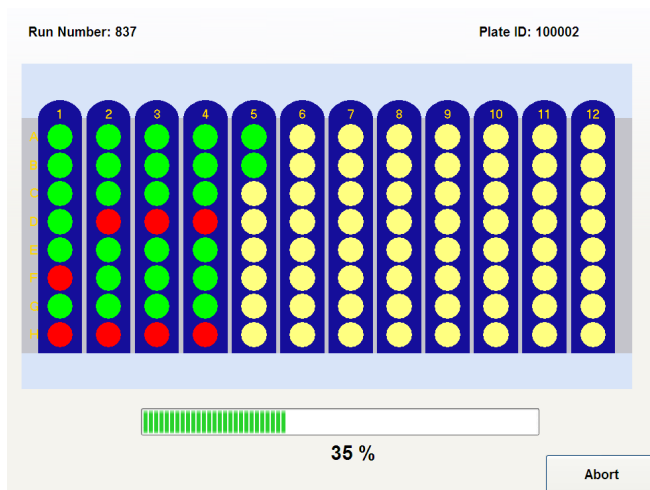


Fig.13: Monitoring the reading process in the CAR's screen. Different colours for different analysis status.

Once finished the reading process, user will be requested again to initialize the opening and closing procedures. (Figs. 12 and 13). Results will be shown once the microplate has been removed from reader and tray is closed.

8.6 SUMMARY REPORT AND RESULTS

SUMMARY REPORT


The CAR generates a final report including following information:

- Date and time
- Position of the sample
- Assay ID
- Sample ID
- Results for each sample

User can look up the final report by selecting "Report View" option. It's possible to print out this run report in PDF format, just by pressing the print option on the upper right side of the screen (Fig. 14).


Genomica SAU
CAR/CAP System


Working list
Run ID: 100002
Date and time: Sat Feb 02 09:59:57 2008



Print

Pos.	Assay ID	Assay	Array ID	Well nr	Sample reference	Result
A1	40204	Papilloma	000000040204	1	1	Genomic DNA control: Passed, Amplification control: No signal, Type 58
B1	40204	Papilloma	000000040204	2	2	Genomic DNA control: Passed, Amplification control: No signal, Type 45
C1	40204	Papilloma	000000040204	3	3	Genomic DNA control: Passed, Amplification control: No signal, Type 18, Type 84
D1	40204	Papilloma	000000040204	4	4	Genomic DNA control: Passed, Amplification control: No signal, Type 33, Type 58
E1	40204	Papilloma	000000040204	5	5	Genomic DNA control: No signal, Amplification control: No signal, Type 6, Type 11, Type 16, Type 51, Type 61, Type 84
F1	40204	Papilloma	000000040204	6	6	Grid alignment failed. The image could not be analysed
G1	40204	Papilloma	000000040204	7	7	Genomic DNA control: No signal, Amplification control: Passed, Type 51
H1	40204	Papilloma	000000040204	8	8	Grid alignment failed. The image could not be analysed
A2	40204	Papilloma	000000040204	9	9	Genomic DNA control: Passed, Amplification control: No signal, Type 33, Type 58
B2	40204	Papilloma	000000040204	10	10	Genomic DNA control: No signal, Amplification control: No signal, Type 6, Type 11, Type 16, Type 51, Type 61, Type 84





Back

Fig. 14: Summary report created by the CAR at the end of the analyzing process..

RESULTS

To visualize a specific sample's result, select the sample/position on the microplate sketch (Fig.15) by pressing twice on it. A new menu, showing working list and results, will appear on the screen (Fig. 16). On this menu, following options are shown:

Export: To export reports and results to an external server, LIMS or USB.

Print: If an external printer is connected via USB port, the selected result's page or all results data pages, including test image, can be printed by operating the "Print" button (See FAQ)

Image: To visualize the image obtained by the CAR (Fig.18).

Raw Data: To consult raw and normalized numerical values obtained in the analysis of the samples.

Virus		
Virus	Result	Controls
Type 6	Negative	Passed
Type 11	Negative	Passed
Type 16	Negative	Passed
Type 18	Negative	Passed
Type 26	Negative	Passed
Type 31	Negative	Passed
Type 33	Negative	Passed
Type 35	Negative	Passed
Type 39	POSITIVE	Passed
Type 40	Negative	Passed
Type 42	Negative	Passed
Type 43	Negative	Passed
Type 44	Negative	Passed
Type 45	Negative	Passed
Type 51	Negative	Passed
Type 52	Negative	Passed

Export
Print
Image
^
V
<< Back Raw Data

Fig.16: Results menu

By selecting the “Image” option, user accesses to the image corresponding to the analyzed sample (Fig. 17 and 18).

Virus		
Virus	Result	Controls
Type 6	Negative	Passed
Type 11	Negative	Passed
Type 16	Negative	Passed
Type 18	Negative	Passed
Type 26	Negative	Passed
Type 31	Negative	Passed
Type 33	Negative	Passed
Type 35	Negative	Passed
Type 39	POSITIVE	Passed
Type 40	Negative	Passed
Type 42	Negative	Passed
Type 43	Negative	Passed
Type 44	Negative	Passed
Type 45	Negative	Passed
Type 51	Negative	Passed
Type 52	Negative	Passed

Export
Print
Image
^
V
<< Back Raw Data

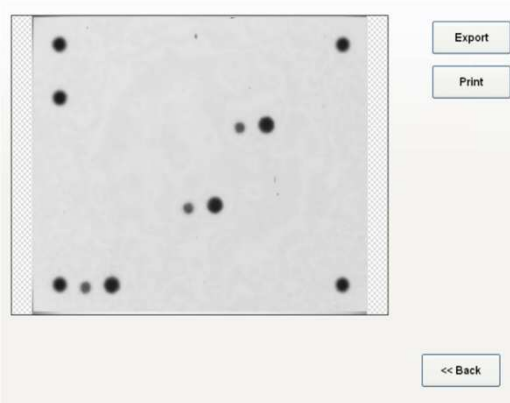


Fig.17: Selecting “Image” option

Fig. 18: Example of a HPV assay performed in strips

By selecting “Raw Data” option, user accesses to the numerical values measured for each virus type (Fig. 19 and 20).

Virus		
Virus	Result	Controls
Type 6	Negative	Passed
Type 11	Negative	Passed
Type 16	Negative	Passed
Type 18	Negative	Passed
Type 26	Negative	Passed
Type 31	Negative	Passed
Type 33	Negative	Passed
Type 35	Negative	Passed
Type 39	POSITIVE	Passed
Type 40	Negative	Passed
Type 42	Negative	Passed
Type 43	Negative	Passed
Type 44	Negative	Passed
Type 45	Negative	Passed
Type 51	Negative	Passed
Type 52	Negative	Passed

Type	Normalized	Normalized	Result
Type 6/11	-0.0078	-0.0081	Negative
Type 16	-0.0012	-0.0013	Negative
Type 18	-0.0011	-0.0011	Negative
Type 26	-0.0032	-0.0033	Negative
Type 31	-0.0064	-0.0066	Negative
Type 33	-0.0086	-0.0089	Negative
Type 35	-0.0053	-0.0055	Negative
Type 39	0.9675	1.0000	POSITIVE
Type 40	-0.0095	-0.0098	Negative
Type 42	-0.0036	-0.0037	Negative
Type 43	-0.0027	-0.0028	Negative
Type 44	-0.0010	-0.0011	Negative
Type 45	-0.0066	-0.0068	Negative
Type 51	-0.0056	-0.0058	Negative
Type 52	-0.0051	-0.0053	Negative
Type 53	-0.0091	-0.0094	Negative
Type 54	-0.0049	-0.0051	Negative

Fig. 19: Selecting “Raw Data” option

Fig.20: Raw data and normalized values

8.7 REANALYZING SAMPLES

After the analysis process, red coloured wells, where the grid alignment has failed, can be reanalyze manually by user.

In the microplate sketch menu (Fig.15), press on a red coloured sample/position to activate the reanalysis main menu window.

Browse the proper Assay among those from the pull down menu (Fig.21) and press “Start Reanalyze”.

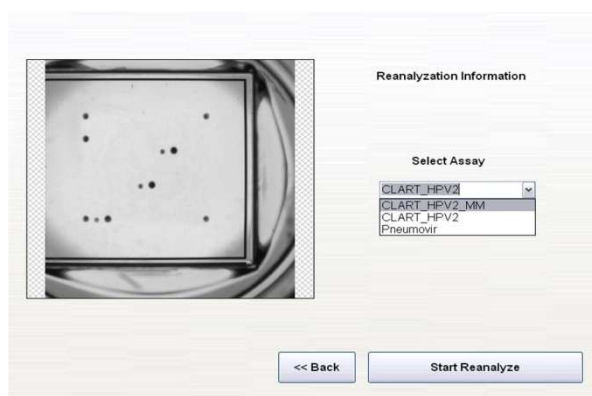


Fig.21: Main menu of the reanalyze option for ASs.

By operating “Start Reanalyze” button, user accedes to the reanalysis application.

The correct grid alignment consist in positioning the upper left marker over the upper left metal or biotin marker and then, adjusting the whole grid size by using the lower right metal or biotin marker. Once the grid is squared properly, according to array's dot micromatrix, press “Reanalyze” button (Fig. 22).

In order to place the upper left marker over its proper position, just press on it and drag it with the finger until both positions match.

Once placed the first marker, user can open or close the grid size by dragging the lower right marker.

Grid movements are limited in order to prevent potential errors when aligning.

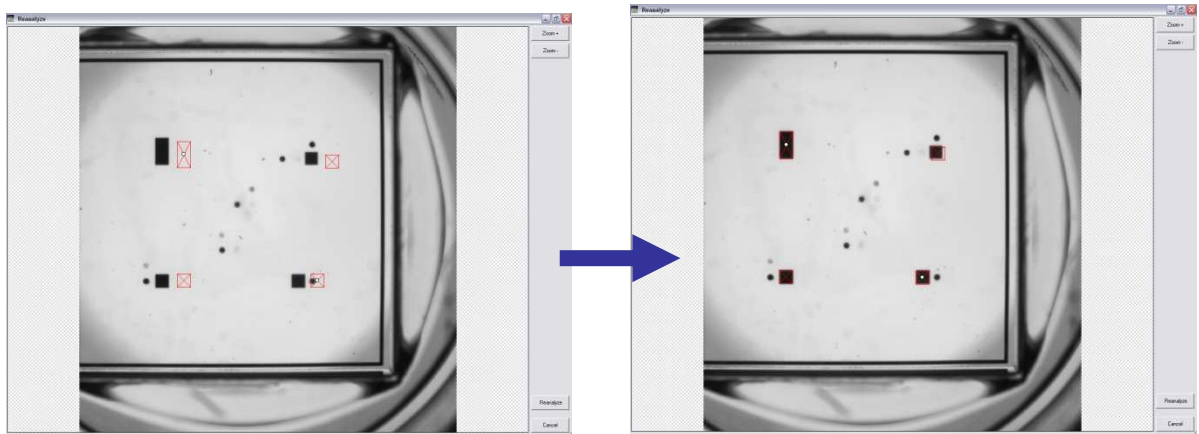


Fig. 22: Placing metal markers according array's matrix.

Zoom tool, placed on the upper right corner of the screen, facilitates the reanalysis process. Scroll the screen to visualize upper and lower markers:

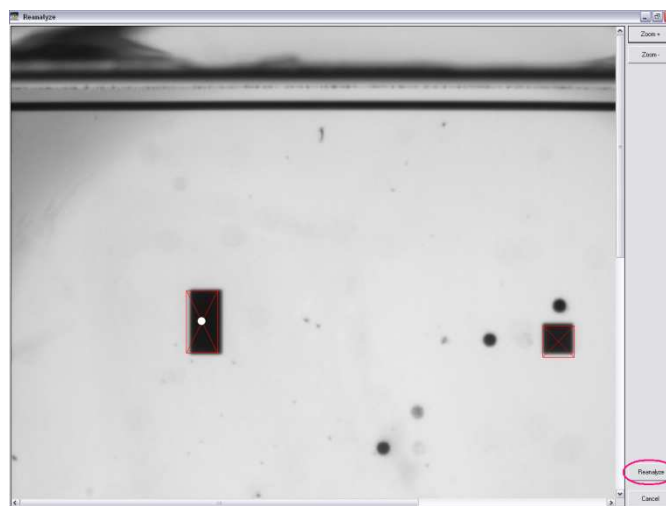


Fig. 23: Manual grid alignment in strips, using metal markers, and zoom tool.

Once the markers are aligned, press “*Reanalyze*” option (Fig 24) and an image with the grid will show the final alignment. If both, biotin and metal markers fits press “*Accept*” (lower- right side). If new adjustments are required, press “*Retry*” (Fig. 24)

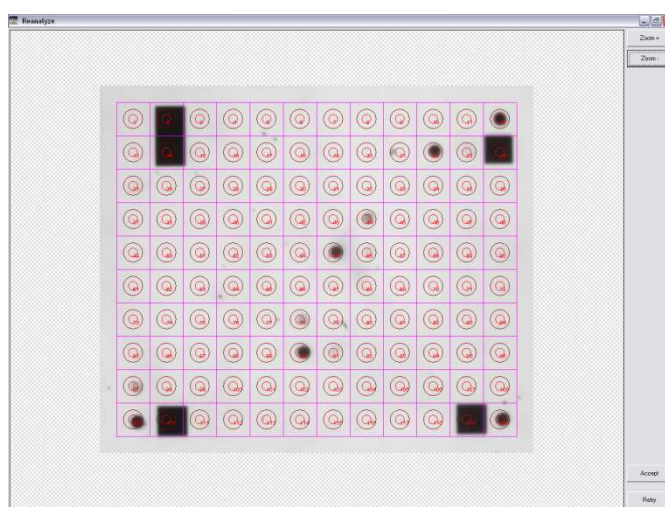


Fig. 24: Grid alignment checking



Manual grid alignment implies user's intervention in the results. If a manual grid alignment is required, an informative message will be shown in the run report.

The image has been reanalyzed by user.

Fig.25: Informative message indicating the manual reanalysis.

9. PERFORMING TUBES

9.1 PLACING ARRAY TUBES ON THE TUBE ADAPTOR

The ATs adaptor is provided by GENOMICA and allows the user to place up to 12 ATs for analyzing.

For the adequate analysis of the tubes, is necessary to place them properly on the adaptor.

Open the AT tube completely and turn back its cap. Hold the tube in this position and introduce both, tube and cap, at the same time into the adaptor as shown in Fig. 25.

Tubes will be read in columns, like the ASs, starting from position A1, which is placed on the blunted square of the tube adaptor (upper left position).

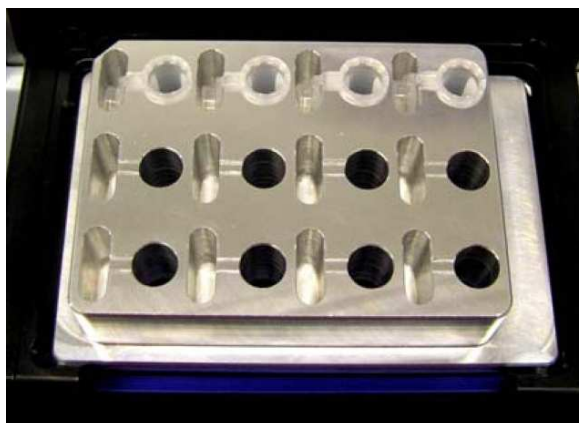


Fig. 25: Placing tubes in the adaptor.

Select “New Analysis” option at the main menu of Genomica’s application (Fig. 26).



Fig. 26: Selecting “New Analysis” at the main menu of GENOMICA’s application.

9.2 CHANGING RACK TYPE

The software establishes the analysis of a whole 96 well plate as default, composed by up to 12 strips..

In case of analyzing ATs (tubes), user should change the *rack type*.

To change the rack type, just select the pull down menu “*Select Rack*” on the top of the screen and choose “*Tube adaptor*” (Fig. 27) instead of “96 well plate”.

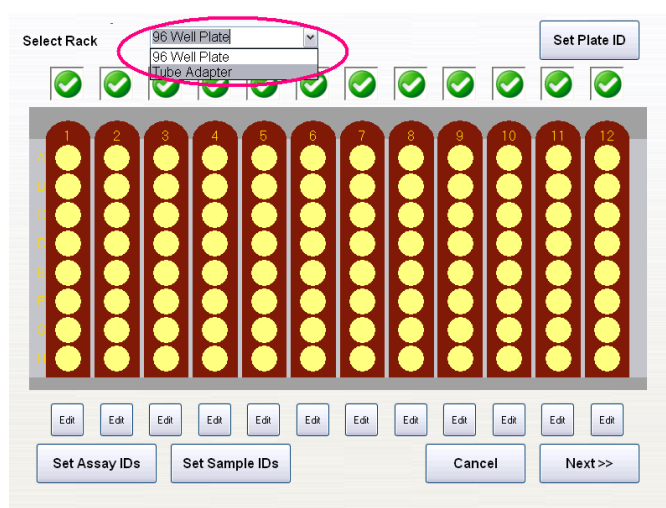


Fig. 27: “Select rack” option at the upper left side of screen.

A new window will pop up showing the tube adaptor sketch (Fig. 28).

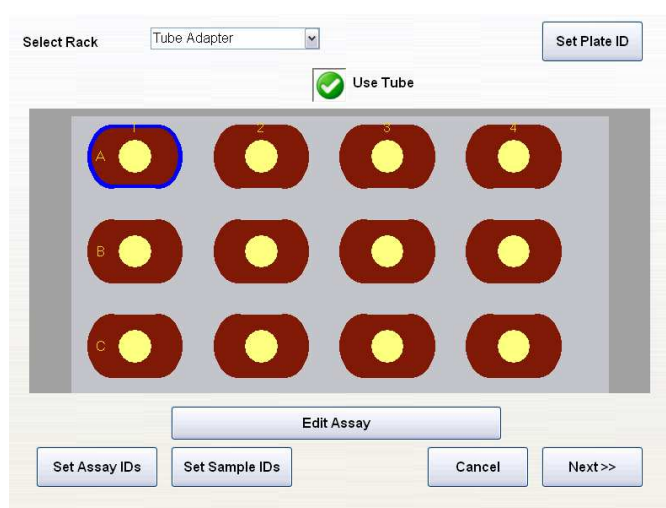


Fig 28: Main menu when reading ATs.

9.3 SETTING SAMPLE'S NUMBER

The number of samples to be analysed can be introduced manually or automatically, by using a barcode reader.

Automatic setting:

For the automatic setting of sample's number, user will need a bar-code reader connected to the CAR (normal USB or PS/2 connection).

Once the adaptor type is changed (see "Changing Rack Type" above) press "Set Assay IDs" option on the lower left side of the screen and all positions will turn unselected (Fig 29 and 30). Read as many barcodes as sample's number you have and tubes will turn to the specific assay's colour combination (Fig. 31).

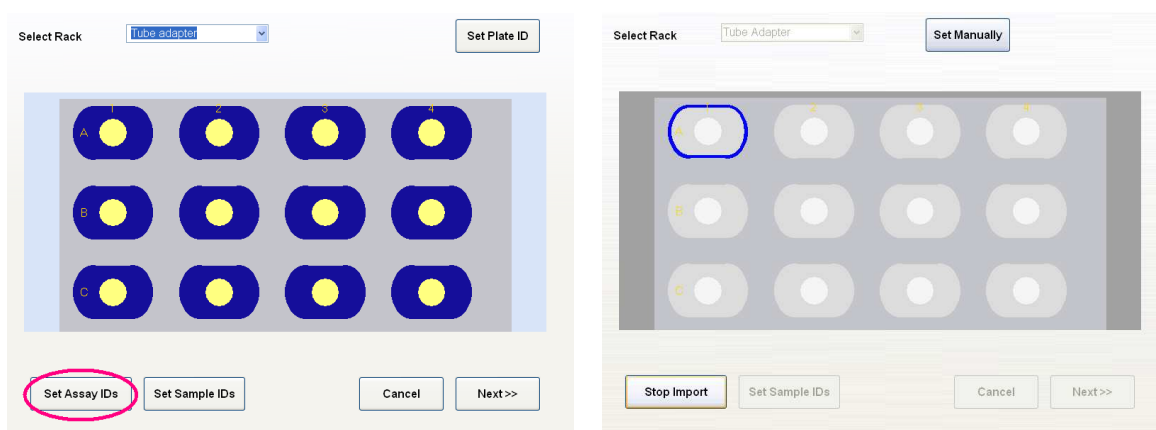


Fig. 29 and 30: Setting Assay IDs automatically

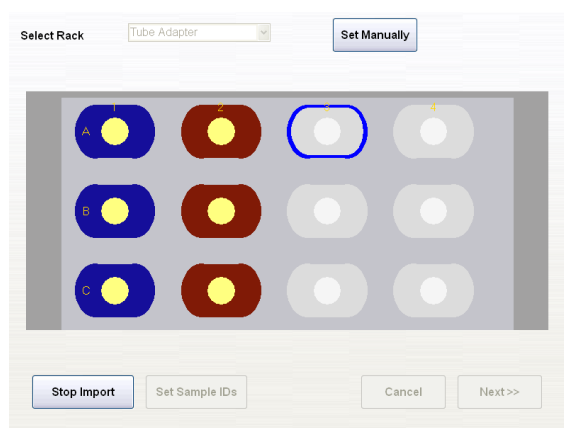




Fig. 31: Different colours combinations for each array type

Manual setting:

In case of analyzing ATs, those tubes that are not going to be processed, should be unselected manually by pressing on  symbol. The no selection of one tube will be indicated with  symbol and the tube position will turn into a light/dark grey combination (Fig. 32).

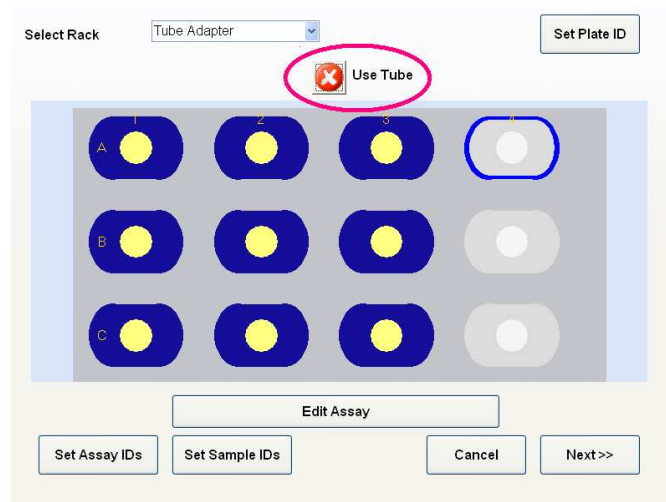


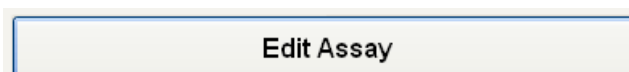
Fig.32: Selected positions appearing in assay's specific combination. Unselected positions turned into light/dark grey combination.

9.4 SETTING ASSAYS IDs

The automatic setting of sample's number, in the previous section, allows the user to skip this step, because assays ID's are already set.

Setting Assays IDs Manually:

Software processes by default the CLART[®] HPV2 Assay. To read or analyse any other GENOMICA'S CLART[®] Assay, select the position (by pressing on it) and then press "Edit Assay" option.



Select the proper assay from those appearing in the pull-down menu as shown (Fig. 33)

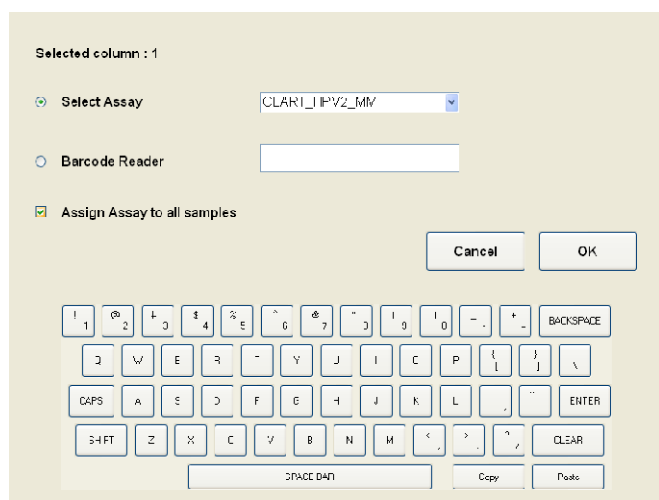


Fig.33: Multiple assays choices.

Software gives also the possibility to select all tubes and edit the Assay ID in all of them at same time. In case of processing all samples with same assay, user should select the “Assign Assay to all samples” on the Edit menu (Fig. 33).

9.5 SETTING SAMPLES IDs

In the same way that in previous sections, setting Samples IDs can be carried out, automatically or manually.



In case of no Samples IDs assignment, software will set automatically a correlative number to each sample, from 1 to 12, starting with the sample placed in position 1A (upper-left).

Setting Samples IDs Automatically



Barcode reader is required to set Samples IDs automatically and barcodes must be set and printed before start setting the IDs.

Set and print barcodes.

Press “Set Samples IDs” option and read the previously assigned barcodes normally. The software associates automatically a barcode to a sample position. Samples with already assigned ID will appear in black (Fig. 34)

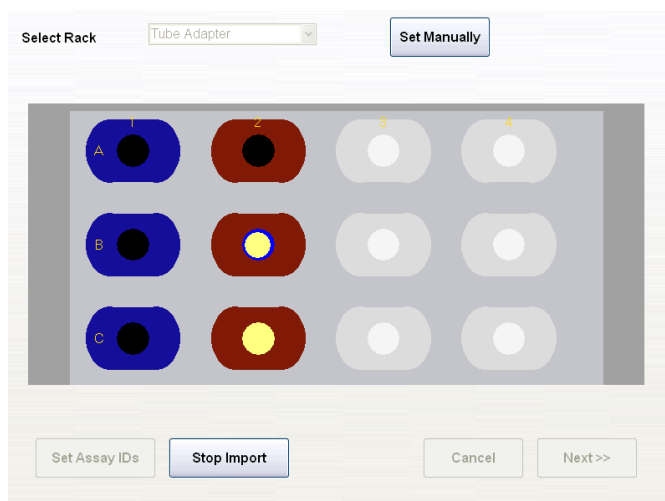


Fig. 34: Setting Samples ID automatically.

Once finished importing IDs, press “Stop Import” to return to the “New Analysis” menu.

Setting Samples IDs Manually

Select each position individually by pressing twice on it. A new window representing a “tactile key board” will pop up on the screen (Fig. 35). Use it to set samples ID manually.

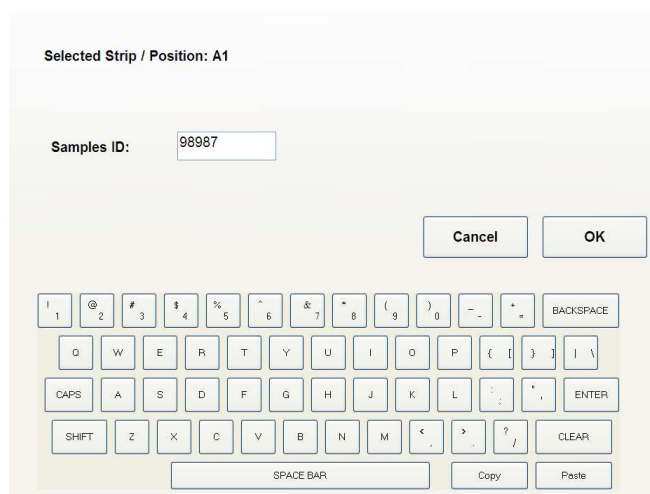


Fig 35: Setting “Samples IDs” manually, using the tactile keyboard.

The user can check the sample position at any time, since it appears at the upper-left side of the screen.

Once finished with the samples’ identification process, press “OK” to go back to the “New Analysis” menu (Fig.29).

Then press “NEXT”.

9.6 WORKING LIST

The working list allows the user to check, whether the already introduced information (samples references, assays...) is correct or not.

To change any data, just press “Back” and repeat the setting process again (see “Set Samples IDs Manually” and “Set Assays ID Manually”).

If all information is correct, press “Start” to carry on with the reading process (Fig. 36).



Pos	Assay ID	Assay	Assay ID	Waiter	Sample reference	Result
A1	40207	CLART HPV2 3.ML	000000040207	-	1	
B1	40207	CLART HPV2 3.ML	000000040207	-	2	
C1	40207	CLART HPV2 3.ML	000000040207	-	3	
A2	40207	CLART HPV2 3.ML	000000040207	-	4	
B2	40207	CLART HPV2 3.ML	000000040207	-	5	
C2	40207	CLART HPV2 3.ML	000000040207	-	6	

Operational Mode: ☐ Process and Reading ☒ Reading

<< Back Start

Fig. 36: Workinglist when performing ATs

9.7 READING PROCESS



Reader's tray opening and closing processes are performed automatically. Do not try to force or move it manually. If necessary, press the emergency button placed on the rear side of the CAR.

User will be requested to initialize the process (Fig. 37), by pressing the “Open Tray” box.

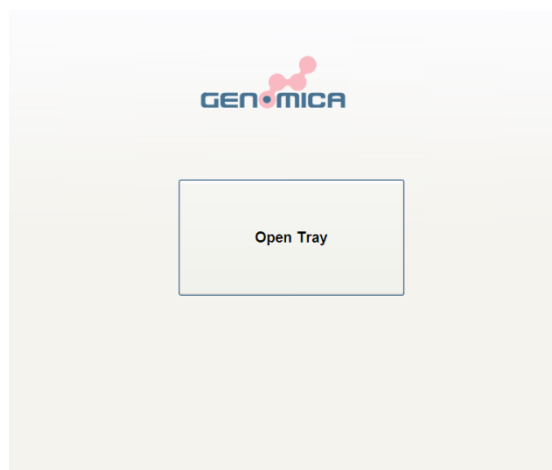


Fig. 37: Message to initialize the reading process.

Once finished the opening process, place the tubes adaptor (ATs) manually in the tray and operate the “Close Tray” box. (Fig.38).



Fig. 38: Message to activate the closing process of reader's tray.

Reading procedure can be monitored in the screen since samples/wells change their colours indicating the reading status (Fig. 39):

- Wells in green: Sample is already analyzed. Grid alignment succeeded.
- Wells in red: Sample may be reanalyzed. Grid alignment failed (see “Sample Reanalyze” section below).
- Wells in yellow: Sample has not been analyzed yet.

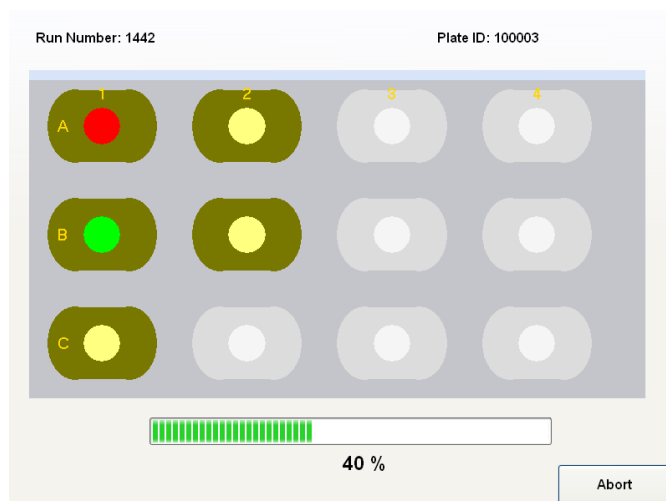


Fig. 39: Reading process

9.8 SUMMARY REPORT AND RESULTS

SUMMARY REPORT

The CAR generates a final report including following information:

- Date and time
- Position of the sample
- Assay ID
- Sample ID
- Results for each sample

User can look up the final report by selecting “Report View” option. From this menu, it is also possible to print it out in PDF format. (Fig. 40).



Fig.40: Summary report created by the CAR at the end of the analyzing process.

RESULTS

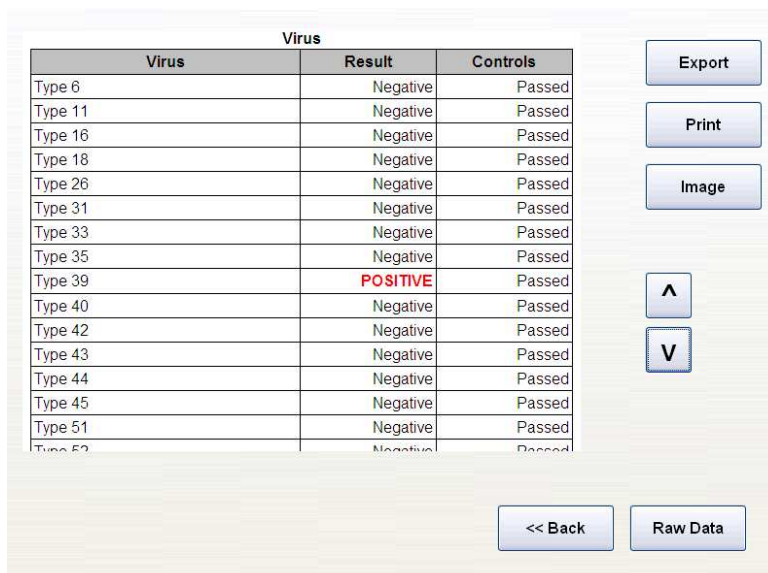
To visualize a specific sample's result, select the sample/position on the microplate sketch screen (Fig.15) by pressing twice on it. A new menu, showing working list and results, will appear on the screen (Fig. 41). On this menu, following options are shown:

Export: To export reports and results to an external server, LIMS or USB.

Print: If an external printer is connected via USB port, the selected result's page or all results data pages, including test image, can be printed by operating the "Print" button (See FAQ)

Image: To visualize the image obtained by the CAR (Fig.18). This image can be exported or printed individually with "print" option

Raw Data: To consult raw and normalized numerical values obtained in the analysis of the samples.



Virus		
Virus	Result	Controls
Type 6	Negative	Passed
Type 11	Negative	Passed
Type 16	Negative	Passed
Type 18	Negative	Passed
Type 26	Negative	Passed
Type 31	Negative	Passed
Type 33	Negative	Passed
Type 35	Negative	Passed
Type 39	POSITIVE	Passed
Type 40	Negative	Passed
Type 42	Negative	Passed
Type 43	Negative	Passed
Type 44	Negative	Passed
Type 45	Negative	Passed
Type 51	Negative	Passed
Type 52	Negative	Passed

Buttons: Export, Print, Image, ^, V, << Back, Raw Data

Fig.41: Results menu

By selecting the "Image" option, user accesses to the image corresponding to the analyzed sample (Fig. 42 and 43)

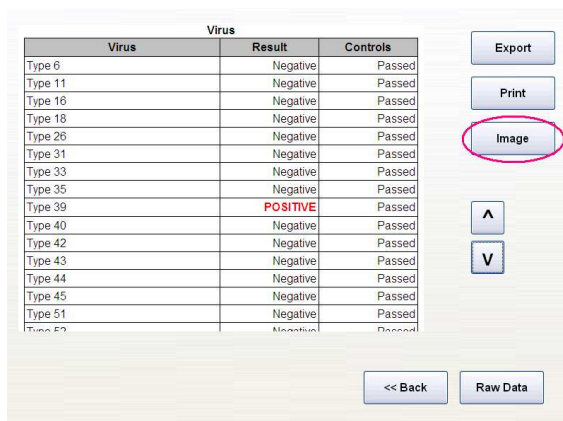


Fig.42: Selecting "Image" option



Fig. 43: Example of a HPV assay performed in strips

By selecting "Raw Data" option, user accesses to the numerical values measured for each virus type (Fig. 44 and 45).

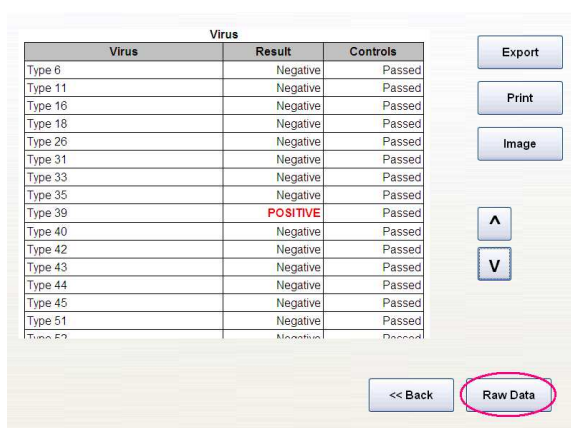


Fig. 44: Selecting "Raw Data" option



Fig.45: Raw data and normalized values

9.9 SAMPLE REANALYSIS

After the analysis process, red coloured wells, where the grid alignment has failed, can be reanalyze manually by the user.

In the tube adaptor sketch menu (Fig.46), press on a red coloured sample/position to activate the reanalysis main menu window (Fig.47)

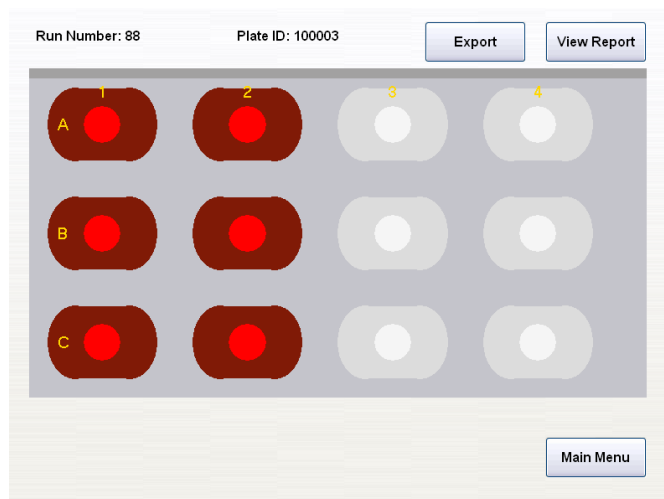


Fig. 46: Grid alignment failed positions when performing Pneumovir Kit.



When performing CLART HPV2 in tubes, alignment must be carried out with the biotin markers instead of Metal Markers. Protocol stays the same

The correct grid alignment consist in positioning the upper left marker over the upper left metal or biotin marker and then, adjusting the whole grid size by using the lower right metal or biotin marker. Once the grid is squared properly, according to array's dot micromatrix, press "Reanalyze" button (Fig 47 and 48).

In order to place the upper left marker over its proper position, just press on it and drag it with the finger until both positions match.

Once placed the first marker, user can open or close the grid size by dragging the lower right marker.

Grid movements are limited in order to prevent potential errors when aligning.

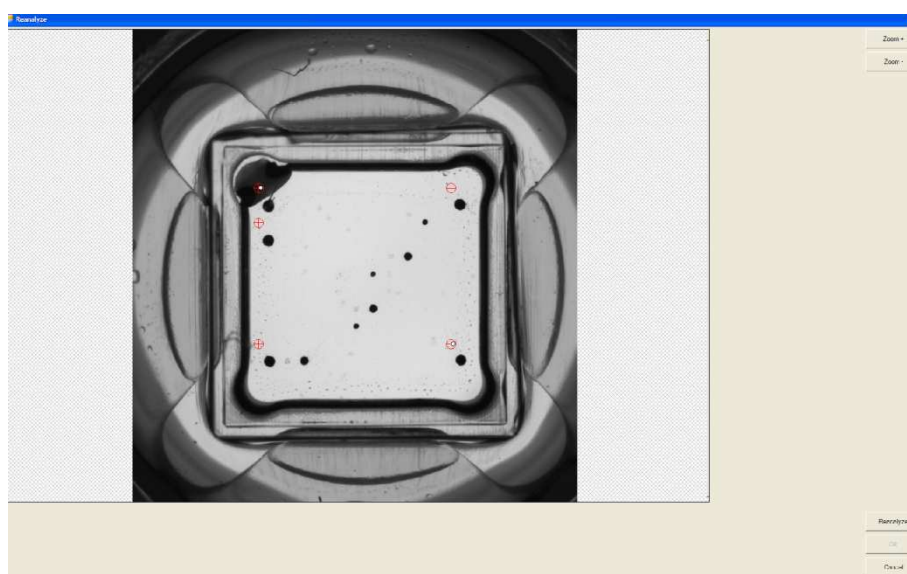


Fig. 47: Manual grid alignment screen.

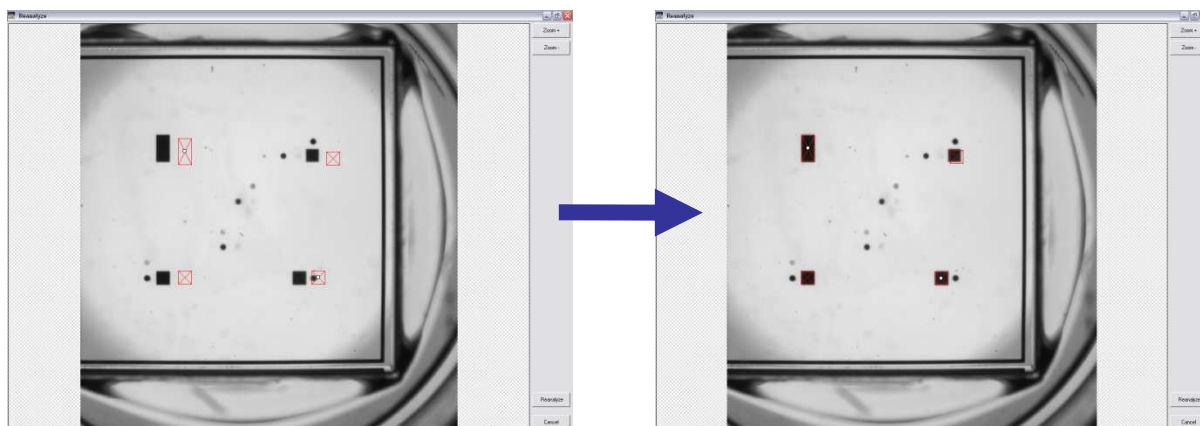


Fig. 48: Aligning the metal markers on the reanalysis menu.

Zoom tool, placed on the upper right corner of the screen, facilitates the reanalysis to user (Fig. 49):

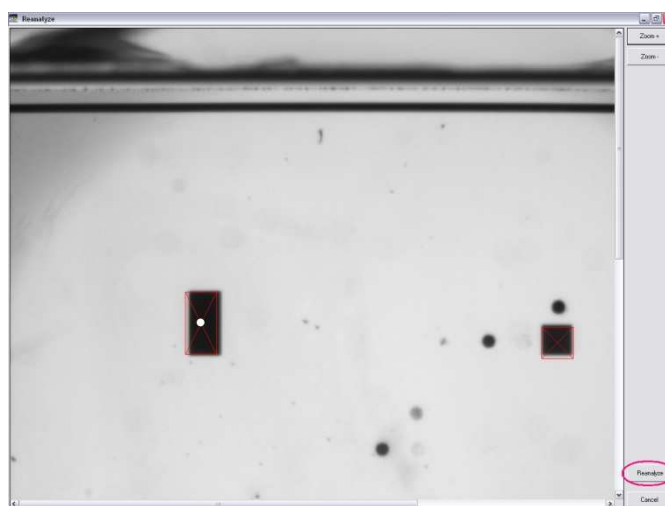


Fig. 49: Manual grid alignment in strips, using metal markers, and zoom tool.

Once the metal markers are aligned, press “*Reanalyse*” option (Fig. 49) and a image with the grid will show the final alignment. If both, dots and metal markers, are well aligned, then press “*Accept*” (lower- right side). If new adjustments are required, press “*Retry*” (Fig. 50)

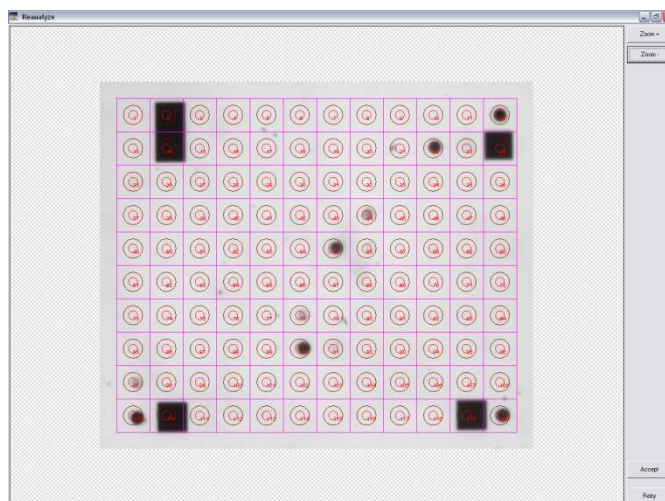


Fig. 50: Grid alignment checking



Manual grid alignment implies user's intervention in the results. If a manual grid alignment is required, an informative message will be shown in the run report.

The image has been reanalysed by user.

Fig.51: Informative message indicating the manual reanalysis.

10. MAIN MENU

10.1 ARCHIVE

By operating the "Archive" option (Fig. 52), user accedes to a result's search engine. Search must be enclosed between two dates, established by the user, a starting date and an ending date (Fig. 53). Set the searching dates required and press "Next".



Fig. 52: "Archive" option on the main menu

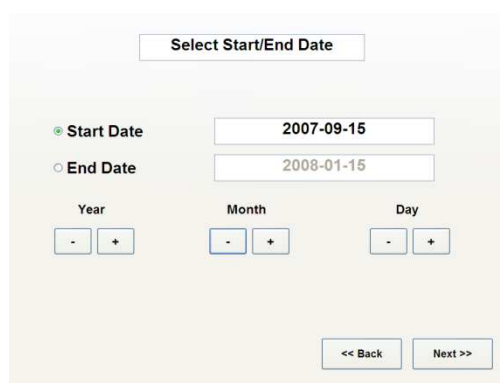


Fig. 53: Menu to set the searching period time.

Search engine will find out all results contained between the two indicated dates. In order to specify the search, three extra searching parameters are available (Fig. 54) in the following menu:

- **Sample ID:** Search samples with same code or ID analyzed during the specified time period.
- **Assay ID:** Search samples processed with same procedure or assay during the specified time.
- **None:** Search for all samples included in the indicated time period.

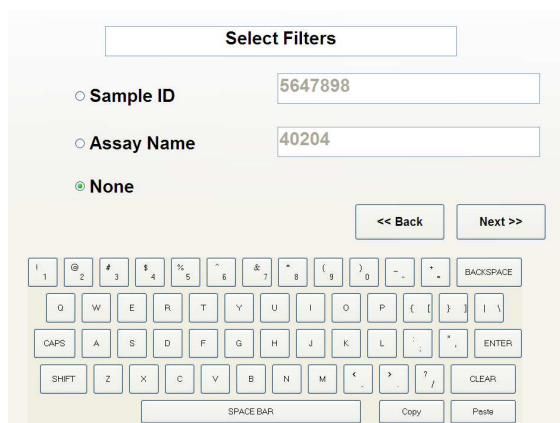


Fig. 54: Available searching parameters.

10.2 SERVICE

By operating the “Service” button (Fig. 55)), user accedes to a submenu containing several reader’s options (Fig. 56):



Fig.55: Selecting “Service” option from the Main Menu.

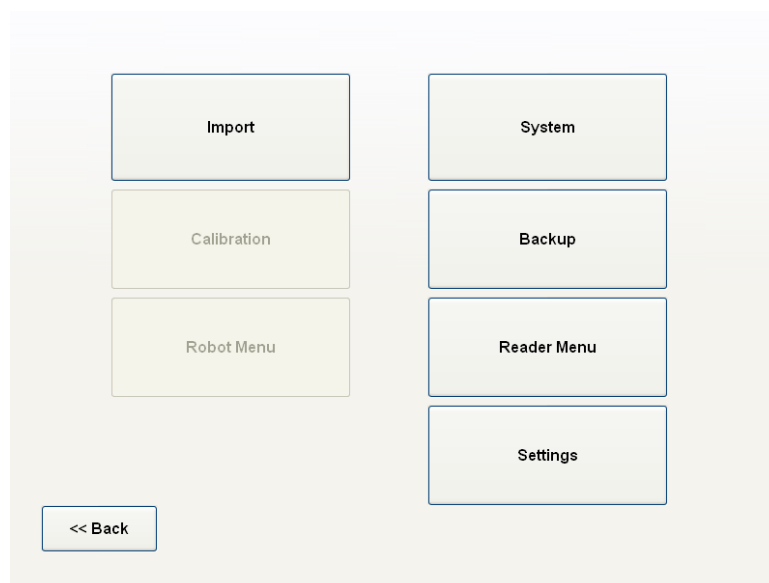


Fig. 56: Multiple options of the "Service" submenu.

10.2.1 IMPORT

This option (Fig. 57) is mainly used for eventual updates in the software version (See section 11.1 for extended explanations).

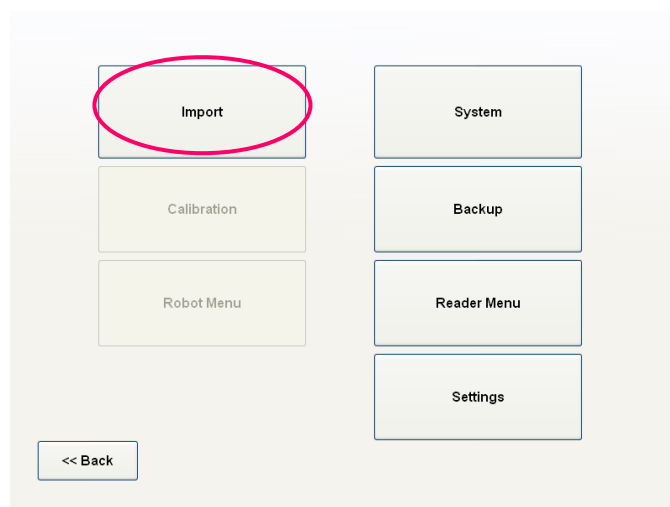


Fig. 57: Selecting "Import" option at the Service Menu.

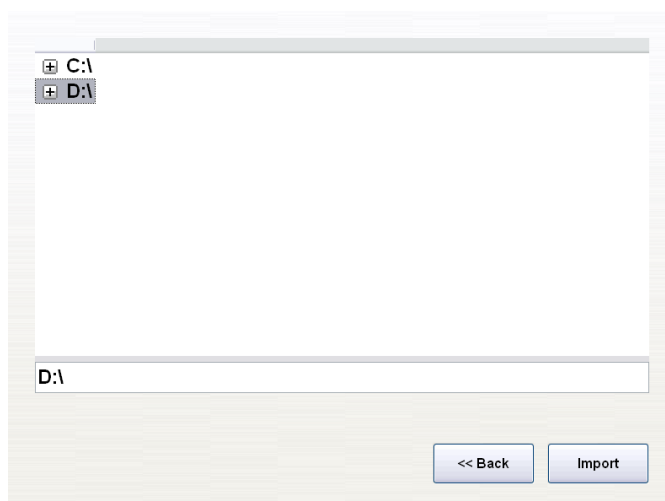


Fig. 58: Multiple importing sources.

10.2.2 SYSTEM

System menu provides the user with information regarding, software, hardware and CAR's serial number (Fig. 59).

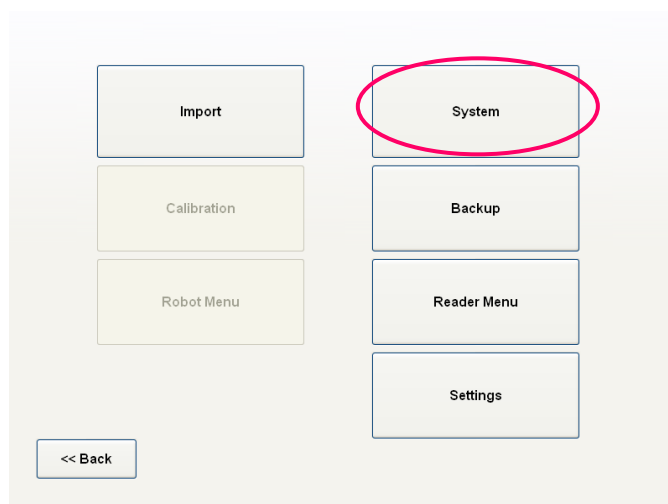


Fig. 59: Selecting System option.

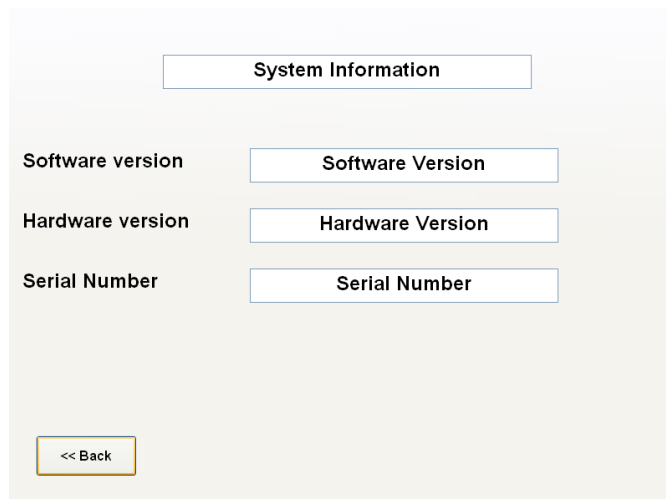


Fig. 60: Information included in "System" menu.

Press "Back" to return to the "Service" menu.

10.2.3 BACKUP/ BACKUP PATH

Reader's storage capacity is limited to 40 Gb, therefore "Backup" option represents a useful tool to save both, data (results, reports...) and CAR's settings.

Moreover, CAR should not be considered as a database where results can be stored. In order to avoid undesirable data lost, backup should be performed regularly.

GENOMICA will not be responsible for the user's data when the technical servicing is provided.

Before saving results with the backup option, user must set the "Backup Path".

SETTING BACKUP PATH

Backup path sets up the target destination for the backup. A remote folder or any other network drive can be specified..

To set the backup path, go to "Settings" option (Fig. 61):

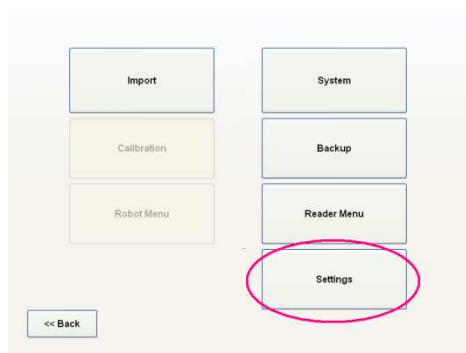


Fig. 62: Selecting “Settings” option at the “Service” menu

Press “...” button:

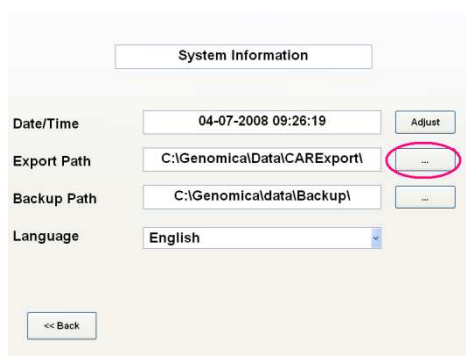


Fig. 63: “...” Box position

Browse directories and external devices connected to the CAR and select the destination (Fig. 64). Press OK when finished.

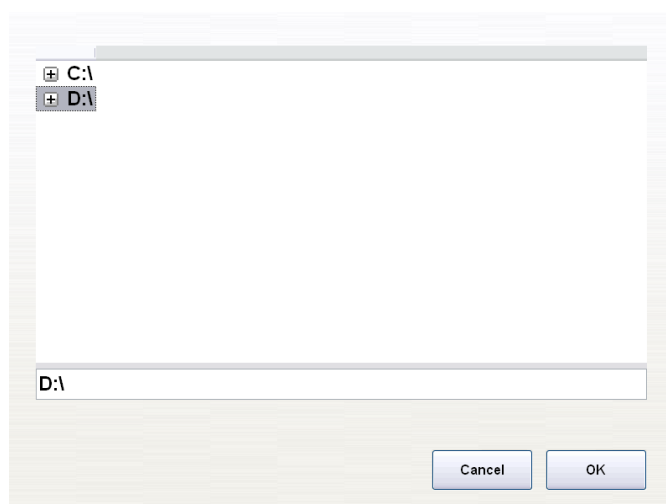


Fig. 64: Selecting directories or external devices.

Once set the backup path, user can backup results by pressing the “Backup” button (Fig. 65):

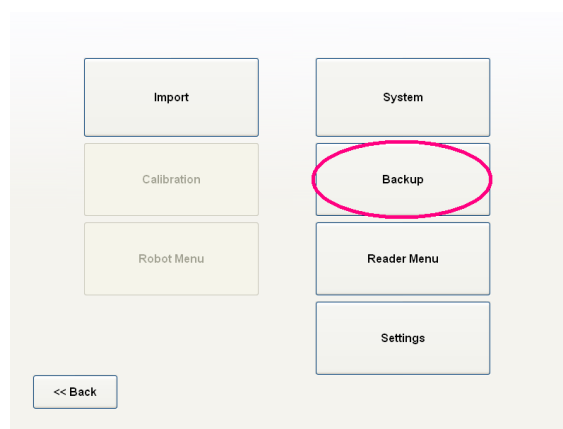


Fig. 65: Selecting “Backup” option.

Next select “Results” (Fig. 66) on the touch screen and then “Next”.

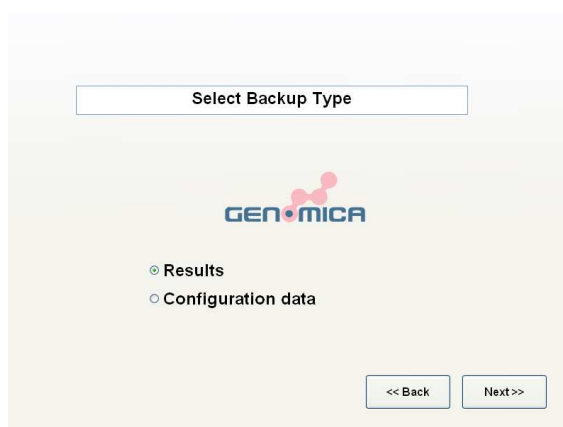
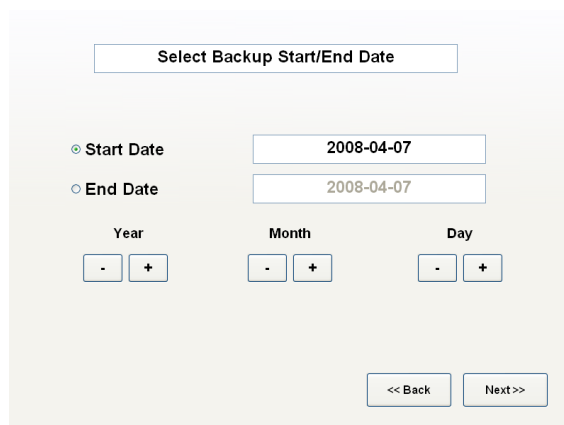


Fig. 66: Selecting “Results” backup.

To speed up data backup, user can introduce a time interval. All results/reports included between the starting date and ending date will be back up (Fig. 67). Then press “Next”.



Select Backup Start/End Date

☒ Start Date 2008-04-07

☐ End Date 2008-04-07

Year Month Day

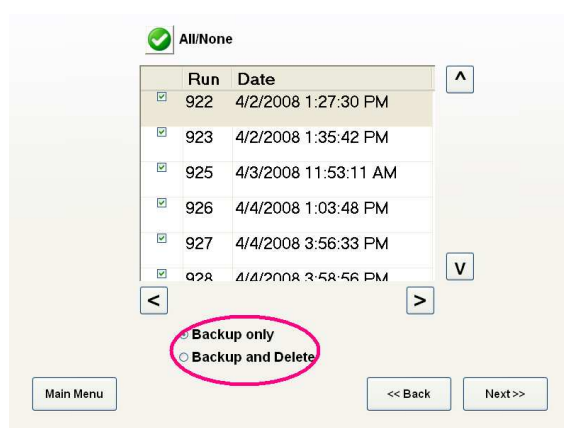
- + - + - +

<< Back Next >>

Fig.67: Introducing dates for the backup

All selected runs are shown on an extra menu (Fig.68).

Both options: “Backup only” or “Backup and Delete” are possible. User decides whether to keep data on the CAR’s reader or erase the already saved results.



✓ All/None

Run	Date
922	4/2/2008 1:27:30 PM
923	4/2/2008 1:35:42 PM
925	4/3/2008 11:53:11 AM
926	4/4/2008 1:03:48 PM
927	4/4/2008 3:56:33 PM
928	4/4/2008 3:58:56 PM

< >

☒ Backup only

☐ Backup and Delete

Main Menu << Back Next >>

Fig. 68: List with selected results and the two backup options.

Press “Next” to finalize the backup process.

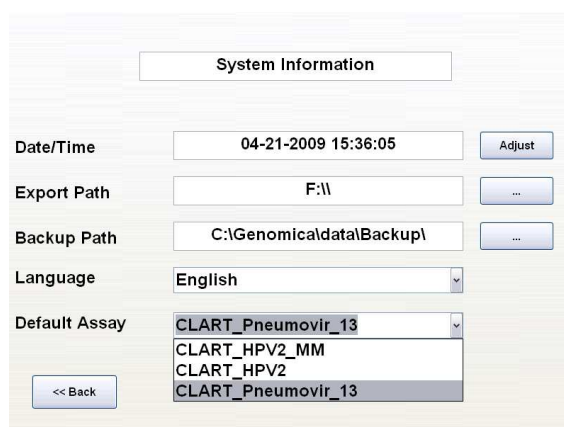


User can change the backup path at any time on the “Settings” menu (See FAQ).

10.2.4 SETTINGS

"Settings" menu (Fig. 69) allows the user to:

- Adjust data/time manually.
- Insert both, data export and backup path.
- Set Language (only English, Spanish, French and Greek are available).
- Set default assay (Pneumovir, HPV2, HPV2 MM, ENTHERPEX)

A screenshot of the "System Information" settings window. It contains several fields and buttons. At the top is a title bar "System Information". Below it are five rows of settings: "Date/Time" with a text box showing "04-21-2009 15:36:05" and an "Adjust" button; "Export Path" with a text box showing "F:\\" and a button with three dots; "Backup Path" with a text box showing "C:\Genomical\data\Backup\" and a button with three dots; "Language" with a dropdown menu showing "English"; and "Default Assay" with a dropdown menu showing "CLART_Pneumovir_13". Below the "Default Assay" dropdown is a list of four options: "CLART_Pneumovir_13", "CLART_HPV2_MM", "CLART_HPV2", and "CLART_Pneumovir_13". At the bottom left is a button labeled "<< Back".

System Information	
Date/Time	04-21-2009 15:36:05 Adjust
Export Path	F:\ ...
Backup Path	C:\Genomical\data\Backup\ ...
Language	English
Default Assay	CLART_Pneumovir_13
<< Back	

Fig. 69:Setting Default Assay

11. FAQ

11.1 SOFTWARE UPDATING

GENOMICA will provide the software necessary to read eventual updates or changes in the kits.



Please take care that the USB used does not contain any virus. GENOMICA is not responsible about virus infections and CAR cleaning will not be consider as a part of the technical support.

Uncompress the zip file we have sent you on your pen-drive.
Check that the pen-drive now contains one folder called "patch" and one or more .bat files with the name of the kit that you want to install. It is important that all of them are in the main directory (e.g. D:\) and not included inside any folder.



Fig. 70

Connect your pen-drive to a free USB connection available on the back of the reader.



Fig. 71:

Go to the “Service” menu.



Fig. 72: Selecting “Service” option.

Press “Import” button.

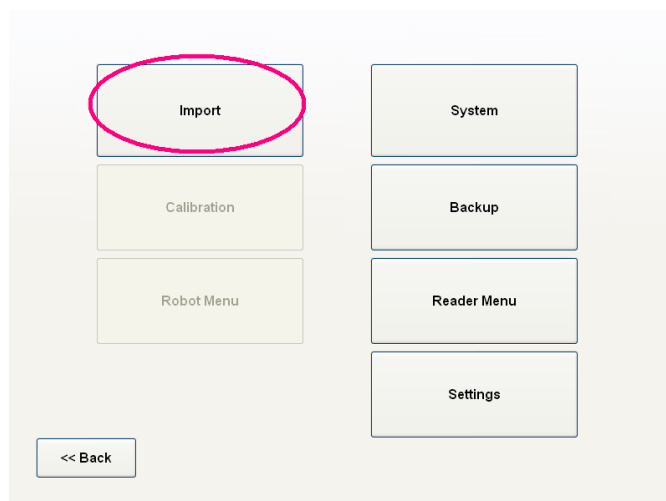


Fig. 73:

In the dialog box you should see both drives: C:\ and D:\ (Corresponding to the USB stick). In some cases, it takes a couple of minutes to the reader to recognize the pen-drive. In those cases, go back to the principal menu and try again in a few seconds.

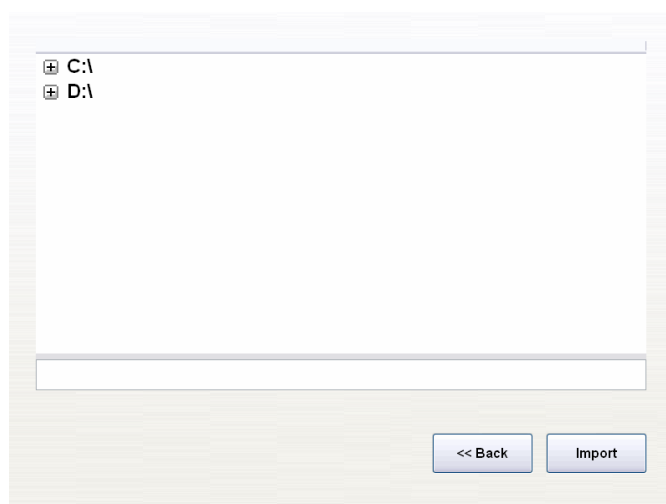


Fig. 74:

Browse the pen-drive contents by pressing on the plus symbol. You can see there the “patch” folder and the kits installation programs (in blue).

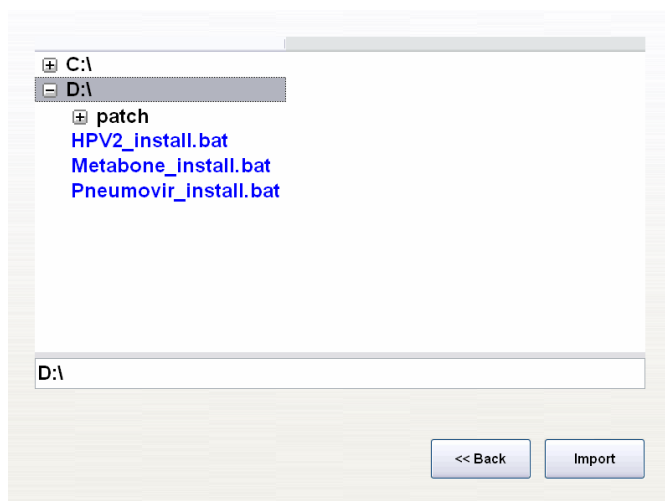


Fig. 75

Select the kit that you want to install and press “Import”.

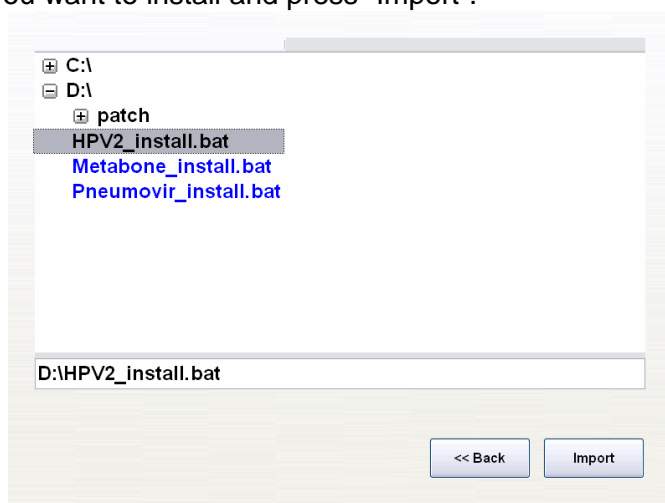
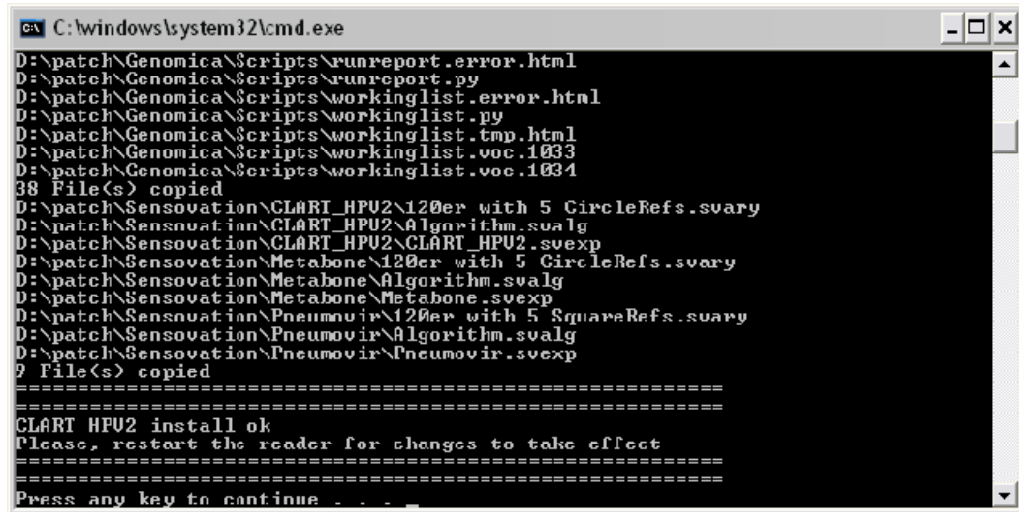


Fig. 76

Installation will be finished when a new window pops up indicating that the process has been completed successfully. Please, close that window by pressing “Cross” button on the top right square. If the installation of more kits is required, repeat the process from step 4. Once finished, get back to the “Main Menu” and reinitialize the reader.

In case that an error occurs during the installation process, please contact the technical service. Do not forget to attach the appearing error messages in your assistance request.



```
C:\windows\system32\cmd.exe
D:\patch\Genomica\Scripts\runreport.error.html
D:\patch\Genomica\Scripts\runreport.py
D:\patch\Genomica\Scripts\workinglist.error.html
D:\patch\Genomica\Scripts\workinglist.py
D:\patch\Genomica\Scripts\workinglist.tmp.html
D:\patch\Genomica\Scripts\workinglist.voc.1033
D:\patch\Genomica\Scripts\workinglist.voc.1034
38 File(s) copied
D:\patch\Sensovation\CLART_HP2\120er with 5 CircleRefs.svary
D:\patch\Sensovation\CLART_HP2\Algorithm.svalg
D:\patch\Sensovation\CLART_HP2\CLART_HP2.svexp
D:\patch\Sensovation\Metabone\120er with 5 CircleRefs.svary
D:\patch\Sensovation\Metabone\Algorithm.svalg
D:\patch\Sensovation\Metabone\Metabone.svexp
D:\patch\Sensovation\Pneumovir\120er with 5 SquareRefs.svary
D:\patch\Sensovation\Pneumovir\Algorithm.svalg
D:\patch\Sensovation\Pneumovir\Pneumovir.svexp
9 File(s) copied
=====
CLART HPV2 install ok
Please, restart the reader for changes to take effect
=====
Press any key to continue . . .
```

Fig. 77

11.2 EXPORTING DATA- EXPORT PATH

User can select where to export all data. Results can be exported to a USB stick, an external hard disk or to the LIMS just by selecting the proper export path.

Steps for exporting results to an USB stick:

Press “Service” option of the Genomica’s main menu:



Fig. 78

Select “Settings” options on the secondary menu:

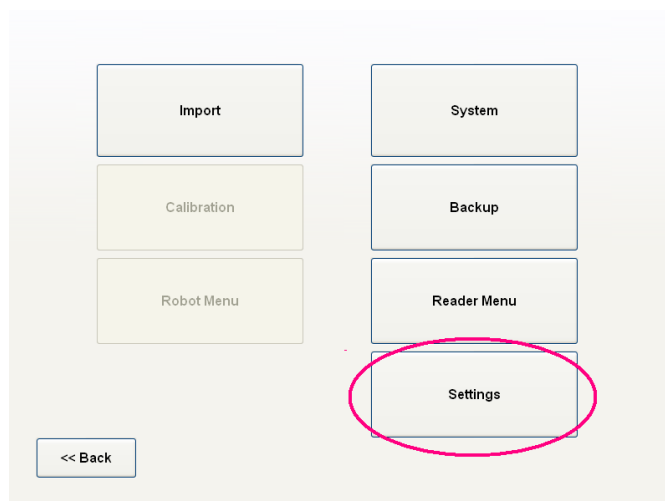


Fig. 79

Browse directories by using the “...” option and select directory which corresponds to the USB stick. Keep in mind that directory assigned to USB may be subjected to changes

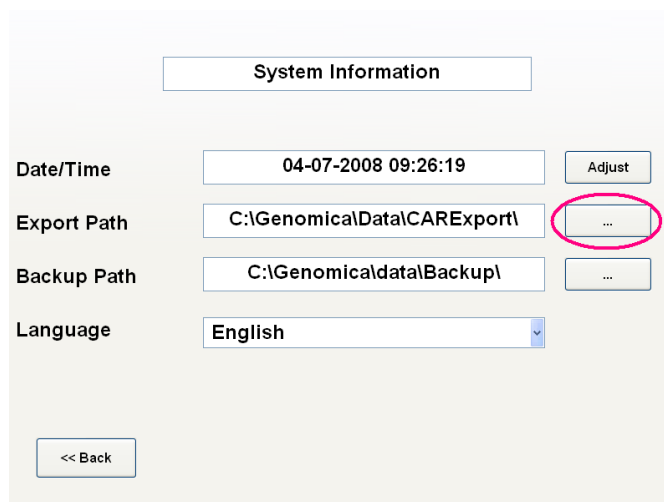


Fig. 80

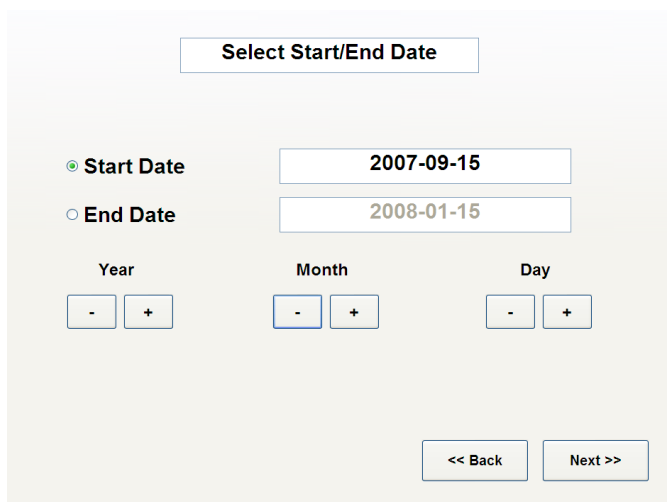
Once selected the export path, all exporting data will be send using the export path. Data exporting should be performed from the “Archive” menu, following the next protocol:

Select “Archive” option from the Main Menu:



Fig. 81

Once selected this option, a searching engine will pop up. On this new window it's possible to restrict the searching time period, inserting a start searching date and end searching date.

The image shows a 'Select Start/End Date' window. At the top is a title bar with the text 'Select Start/End Date'. Below it are two radio buttons: 'Start Date' (selected) and 'End Date'. To the right of these are two date input fields: '2007-09-15' and '2008-01-15'. Below the date fields are three groups of controls for 'Year', 'Month', and 'Day'. Each group consists of a '-' button and a '+' button. At the bottom right are two buttons: '<< Back' and 'Next >>'.

Select Start/End Date		
<input checked="" type="radio"/> Start Date	2007-09-15	
<input type="radio"/> End Date	2008-01-15	
Year	Month	Day
- +	- +	- +
<< Back Next >>		

Fig. 82

Search engine will find out all results contained between the two indicated dates. In order to specify the search, three extra searching parameters are available:

- **Sample ID:** Search for samples with same code or ID analyzed during the specified time period..
- **Assay ID:** Search samples processed with same procedure or assay during the specified time.
- **None:** Search all samples included in the indicated time period.

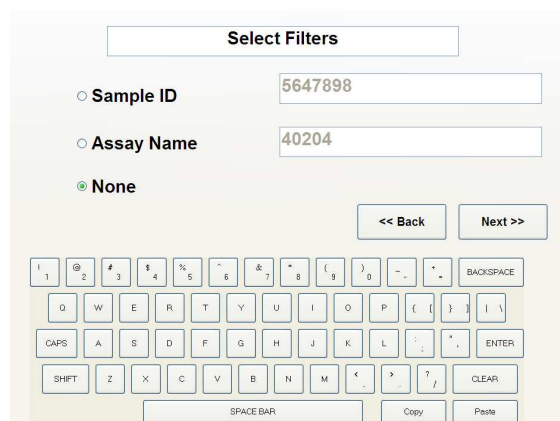
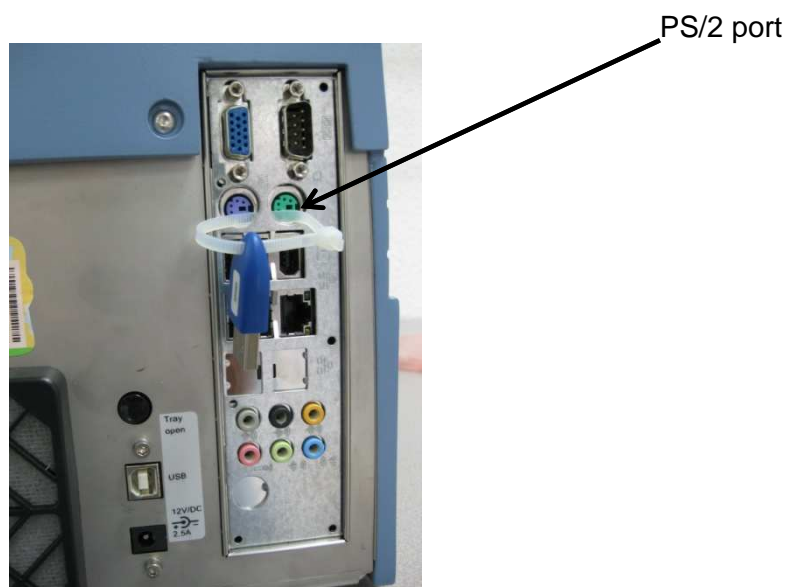


Fig. 83

Press “Next” to continue.

11.3 BARCODE READER: CHARACTERISTICS AND INSTALLATION

A simple barcode reader will be advisable since installation process is done automatically by the device just by connecting it at the rear PS/2 or USB port.



Barcode Reader specifications (recommendation):

Scanner	Type	Barcode Scanner
	Scanning Element	CCD
	Light Source	660 nm red visible LED
	Decode Capability	UPC-E, UPC-A, EAN-8, EAN 13, JAN, UCC / EAN 128, ISBN, ISSN, Code 39, Code 93, Code 128, Interleave 2 of 5, MSI, Industrial 2 of 5, Matrix 2 of 5, IATA 2 of 5, China Post Code, Code 32, Codabar, ABC Codabar, CX Codabar, Code 11, UK Plessey, Telepen
	Scanning Speed	45 scans per second
	Decode Speed	100 scans per second
Barcode	Max Working Distance	1"
Connectivity	Interface	PS/2
Power Requirements	Voltage	+5 VDC
	Consumption	Decoded: Standby <30 mA, Working <95 mA
Operating Conditions	Temperature	0 to 50 Degrees C
	Humidity	20% to 80% non condensing

11.4 DONGLE. WHAT IT IS AND ITS USE

Dongle is the name of an electronic key with USB appearance. Depending on the CAR's serial number, Dongle can be found, whether the rear side of the CAR, attached with a bridge, or internally (not visible)

When the Dongle is placed outside, in the rear side, just insert the Dongle in a free USB port in order to let the CAR work. Without the Dongle, CAR will start a maximum of 30 times before stop working.

When the Dongle is placed inside the CAR, it will be indicated in a sticker, at the rear side.

Dongle works as a software license. In case of Dongle failure, GENOMICA application won't be launch and CAR won't be operative.

Please, do not remove the Dongle from the CAR unless for shipping it to technical service.



12. APPENDIX 1

ABBREVIATIONS USED

AS: Array Strips

AT: Array Tube

CAR: Clinical Array Reader

Fig. : Figure

PCR: Polymerase Chain Reaction

PDF: Portable Document Format generate by Adobe.

LIMS: Laboratory Information Management System

CLART: Clinical Array Technology

13. APPENDIX 2

SYMBOLS AND WARNING SIGNS



Yellow triangles indicate general warnings. Warnings include the risk injury to the user, or serious equipment damage.



This symbol indicates severe injury to the instrument operator due to interference of the user with moving parts of the *CAR*.



This symbol indicates the risk of loss of life or severe injury to the user due to high voltage.



This symbol indicates important information about the *CAR* and analyzing process. Please read these instructions carefully.



This symbol indicates important information regarding ATs analyse.